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> **Callipeltin I** N H H $N \rightarrow N$ H O O O NH HN N H_2N $\begin{bmatrix} N \\ N \end{bmatrix}$ O O O OH OH OH $\begin{array}{ccc} \nabla H & \mathbf{O} & \mathbf{O} & \rightarrow \\ \nabla \mathbf{H} & \mathbf{O} & \mathbf{H} & \mathbf{O} \end{array} \begin{array}{ccc} \nabla \mathbf{H} & \mathbf{O} & \mathbf{O} & \mathbf{H} \\ \nabla \mathbf{H} & \mathbf{H} & \mathbf{O} & \mathbf{H} \end{array}$ <code>N \searrow он</code> O O NH $_{\rm H_2}$ N $^{\sim}$ NH NH_2

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$$
R^{1}C \equiv CH + (CH_{2}O)_{n} + HNR^{2}R^{3} \xrightarrow{\text{Cul/Al}_{2}O_{3}} R^{1}C \equiv CCH_{2}NR^{2}R^{3}
$$

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> OH OH R

N

 $N_{\rm N}$

O

N

 $(+/-)$ $(+/-)$

NHMs OMs

X

Gilles Gosselin, Ludovic Griffe, Jean-Christophe Meillon* and Richard Storer

5 steps

 $O_{\scriptscriptstyle\diagdown\hspace{0.3pt}}$ $O_{\scriptscriptstyle\diagdown\hspace{0.3pt}}$

O

O R

R_O

r Y o

R'

 H_O

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 $NADP^+$ D-Glucose

OH OH

HO

or \bigwedge $\begin{matrix} N \\ N \end{matrix}$ $\begin{matrix} Y \end{matrix}$

HO

R

O

 $I-Ms$

 $N_{\rm N}$

N

X

N

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⁽i) Cyanuric chloride, NEt₃, DCM, 0 \degree C to rt, 8 h (85-98%) R = aryl, alkyl, benzyl, $o\text{-}C_fH_a\text{-}NHCOCH=CHCOOH$; R' = H, Ph; R'' = H, Me; R'R'' = \mathbb{R}

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N $Ac\Omega$ AcO O O O $R¹$ (*L*)-Tartaric acid $\implies \qquad \searrow N$ TMSO TMSO O O R $\searrow N$ **TMS** TMSO_(ii) O OH Ń∙R R^1 N $\Delta \cap$ AcO O OAc R R^1 R = R^1 = Ph, *i*-Pr, vinyl, c-hexyl, Me, Ph O O $R¹$ MgX Ac₂O
DMAP BF_3E_2O various solvents and reaction conditions up to 93% de

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PhCN

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A synthon approach to spiro compounds

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Abbreviations: BINAP, 2,2'-bis(diphenylphospospiro)-1,1'-binaphthalene; Boc, t-butoxycarbonyl; CTAB, cetyltrimethylammonium bromide; DEAD, diethylazodicarboxylate; 4-DMAP, 4-(dimethylamino)pyridine; Fmoc, 9-fluorenylmethoxycarbonyl; HMPA, hexamethylphosphoric amide; HMPT, hexamethylphosphorous triamide; KHDMS, potassium hexamethyldisilazide; LAH, lithium aluminium hydride; LDA, lithium diisopropylamide; LiHDMS, lithium hexamethyldisilazide; m-CPBA, m-chloroperbenzoic acid; NaHMDS, sodium hexamethyldisilazane; NBS, N-bromosucinimide; NCS, Nchlorosuccinimide; NMO, N-methylmorpholine N-oxide; PCC, pyridinium chlorochromate; PDC, pyridinium dichromate; Ph, phenyl; PhH, benzene; PhMe, toluene; PLG, L-propyl-leucyl-glycinamide; PMP, 1,2,2,6,6-pentamethylpiperidine; PPTS, pyridinium toluene-4-sulphonate; rt, room temperature;
TFA, trifluoro acetic acid; THF, tertrahydrafuran; TMEDA, *N,N,N',N'*-tetr TPAP, tetrapropylammonium perruthenate; p-TSA, p-toluene sulphonic acid; Z, benzyloxycarbonyl.

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1. Introduction

In 1900, Bayer created the first spiran described as a bicyclic hydrocarbon connected by a single carbon. The term spirocyclanes was used to describe the family of such hydocarbon. Due to the tetrahedral nature of the spirolinked carbon, the ring planes are nearly perpendicular to each other.

Spiro compounds having cyclic structures fused at a central carbon are of recent interest due to their interesting conformational features and their structural implications on biological systems. The asymmetric characteristic of the molecule due to the chiral spiro carbon is one of the important criteria of the biological activities. The retention of neurotoxic properties of perhydrohistrionicotoxin (1), an analogue of a natural product 2, is clear evidence of the role of the spiro carbon in steering the biological activity.^{[1](#page-52-0)}

2. Biological activities

The spiro functionality has been known for a long time to be present in phytochemicals either in alkaloids, lactones or terpenoids. The spirocyclic alkaloid $(-)$ -histrionicotoxin (2), isolated from skin extracts of the poison dart frog, Dendrobats histrionius, found in Columbia, is a very potent nicotinic receptor antagonist.^{[9](#page-52-0)} Spiroketals are reported to be the sub-units of many naturally occurring substances of biological interest such as insect pheromones, antifeedants and polyether antibiotics.^{[10](#page-52-0)} A series of spiroketals $(3-6)$ have been isolated from Chrysanthemum coronanium, a common vegetable of South China.^{[11](#page-52-0)} Some of these compounds are found to have antifeeding activity towards silkworm^{[12](#page-52-0)} and spasmolytic and antiphlogistic activity.^{[13,14](#page-52-0)} Unsaturated spiroacetals such as 1,6-dioxaspiro[4.4]nona-3,8-diene^{[15](#page-52-0)} and 1,6-dioxaspiro^[4.5]decane^{[16](#page-52-0)} have also been

The spiro [pyrrolidin-3,3 $'$ -indole] ring system is a recurring structural motif in a number of natural products such as vinblastine and vincristine, that function as cytostatics and are of prime importance in cancer chemotherapy.[17](#page-52-0) The derivatives of spiro-oxindole find very wide biological application as antimicrobial, antitumour, and antibiotic agents, and inhibitors of human NK-1 receptor etc.¹⁸⁻²⁰ Horsfiline (7), an oxindole alkaloid containing a spiro- [indole-pyrrolidone] nucleus, has been isolated by Bodo and co-workers^{[21](#page-52-0)} from *Horsfieldia superba*, a tree from Malaysia, the extracts of which are commonly employed in local medicine.

The saponaceolides (A–D), 8 are found to possess antitumour activity in 60 human cancer cell lines. 22 22 22 Each of these compounds contains a unit of tricyclic trioxaspiroketal.

The presence of the sterically constrained spiro structure in various natural products also adds to the interest in the investigations of spiro compounds.^{[2](#page-52-0)} Spiro compounds represent an important class of naturally occurring substances characterised by their highly pronounced biological properties. $3-5$

In the arena of photochromism, spiro compounds, due to their steric constraints, equilibrate with the corresponding non-spiro analogue and exhibit various photochemical phenomena. The discovery of the photochromic reactions of spiropyrans^{[6](#page-52-0)} (Scheme 1) during 1952 initiated the work in the area of photochemical erasable memory.^{[7](#page-52-0)}

Some more related applications based on the equilibrium are self-development photography, actinometry, displays, filters and lenses of variable optical density etc.

The alkaloids containing a spiro[indole-pyrrolidine] nucleus are cell-cycle-specific cytostatic agents that arrest mitosis and metaphase by acting as spindle poisons. They are also found to be useful in cancer chemotherapy.^{[23](#page-52-0)}

Some spiroacetals (9) and (10) show strong cytotoxic activity against human cancer cells.^{[24](#page-52-0)} These compounds are hybrid natural products made from estrone (11) and a highly biologically active mycotoxin, talaromycin B (12) .^{[25](#page-52-0)}

idene-spiro-hydantoin analogue (16) of hydantocidin was shown to be the most efficient inhibitor of muscle glycogen phosphorylase B known to date. The thio-analogue (17) is a potent inhibitor of glycogen phosphorylase B and glycogen phosphorylase A, not only from muscle, but also of liver \arcsin^{30} \arcsin^{30} \arcsin^{30}

A research group at Merck 31 has developed a neuropeptide antagonist, which exhibits antidepressant activity.

 R_7 and R_8 = H, halo, alkyl, carboxy, alkoxy methyl, carbamoyl, etc.

Napalilactone $(13)^{26}$ $(13)^{26}$ $(13)^{26}$ and pathylactone A $(14)^{27}$ $(14)^{27}$ $(14)^{27}$ are novel norsesquiterpenoid spirolactones isolated from marine sources and $\overline{14}$ was reported to be a Ca^{2+} antagonist.^{[28](#page-52-0)} $(+)$ -Hydantocidin (15), which contains a unique spironucleoside structure, possesses potent herbicidal and plant-growth regulatory activity.^{[29](#page-52-0)} The D-glucopyranosylMaligres and coworkers^{[32](#page-52-0)} have synthesised a non-peptidal $(-)$ -spirobyclic NK-1 (18) receptor antagonist, which is required for their clinical programme.

The azaspiro compounds (19) are reported^{[33](#page-52-0)} to be tachykinin antagonists and are of particular use in the treatment of depression, anxiety, pain, inflammation, migraine, emesis or postherpetic neuralgia.

Some spiroheterocycles, benzopyrans (20) and (21), are aldose reductase inhibitors, which are found to be useful as antidiabetics.[34](#page-52-0) Several potent reductase inhibitors based on spirosuccinimide, spiropyridazine and spiroazetidine have been reported for the prevention of secondary complications of diabetes.[35](#page-52-0)

Spirobicyclic lactam peptidomimetics^{[36](#page-53-0)} (22) and (23) of L-prolyl-leucyl-glycinamide (PLG) were found to exhibit a pharmacological profile similar to PLG (24) in terms of their $\int_0^3 H$]-spiroperidol/N-propylnorapomorphine D₂ receptor competition binding $\arccos 37,38$ $\arccos 37,38$ and the 6-hydroxydopamine-lesioned animal model of Parkinson's disease. Both (22) and (23) produced a greater shift from the low-affinity state to the high-affinity state of the dopamine $D₂$ receptor than PLG. Spirocyclic quinuclidines (25) containing spirofused indoles are important for muscarinic receptor binding.^{[39](#page-53-0)} Substituted quinuclidines exhibit selective muscarinic receptor binding properties.⁴

 $R=H, C_5H_9, C_6H_5,$ i-Pr, 4,4-diethoxy butyl Spirolide R_1 R_2

Spirolides A–D (26–29) and 13-desmethyl spirolide C (30) are toxic in a mouse bioassay for lipophilic toxins.^{[41–44](#page-53-0)} These compounds were first found in the extract of shellfish from aquaculture sites and, subsequently, they were shown to be produced by dianoflagellates such as Alexandrium ostenofeldii. All the compounds (26–29) contain an unusual seven-membered spiro-linked cyclic imine moiety with the same atomic connectivity as found in pinnatoxins A–D (31–34), which are potent calcium channel activators and are responsible for outbreaks of shellfish poisoning in China and Japan.[45](#page-53-0)

Cordi and co-workers 46 have reported some spiro-imidazoline compounds endowed with α -adrenergic agonist activities. They have also reported that (R,\bar{S}) -spiro $(1,3-)$ diazacyclopent-1-ene)-5,2'- $(7'$ -methyl-1',2',3',4'-tetrahydronaphthalene) (35) is a prototype of venospecific constrictors as it is able to constrict the saphenous vein of dogs without noticeable effects on the mean arterial pressure.[47](#page-53-0)

Synthetic compounds with a 2-spiropiperidine moiety also possess interesting pharmacological activities.

Spiro[piperidine-2,2'-adamantane] (36) proved to be active against influenza viruses 48 and the spiro[piperidine- $2,3'(2'H)$ -benzopyran] (37) showed a significant and selective affinity for the $5-\text{HT}_{1\text{A}}$ receptors^{[49](#page-53-0)} and, therefore, serves as an anxiolytic agent.

oxidation and reduction, offering the possibility of readout and writing in either optical or electrical mode. The photoisomerisation of the azo group, however, has been shown to have the disadvantages of relatively low photosensitivity and a limited wavelength region for the

Interesting biological activities have been found in the piperidine alkaloids, (\pm) -pandamarine (38) and (-)pandamarilactone (39), which contain an azaspiro[4.5]- decane structural unit.^{[50](#page-53-0)} The alkaloids (38) and (39) are isolated from *Pandanus sp.*^{[51](#page-53-0)} Pinnaic acid (40) and tauropinnaic acid (41), isolated from *Pinna muricata*,^{[52](#page-53-0)} are found to exhibit inhibitory activity against a cytosolic 85 kD phospholipase (cPLA₂).^{[53](#page-53-0)} Halichlorine (42), isolated from the marine sponge Halichondria okadi, is an inhibitor of the vascular cell adhesion molecule.^{[54](#page-53-0)}

coloured form.^{[61](#page-53-0)} These difficulties hinder the development of azo compounds for practical molecular switching devices. In contrast, a feature making spiropyrans one of the most widely studied classes of photochromic compounds is the intense absorption of the coloured form in the visible region. This is of great importance for practical applications of spiropyrans in display and photochromic memory systems. 62 Spiro compounds, due to their equilibrating cyclic and acyclic structures, are well established as photochromic materials.⁶³⁻⁶⁷

The spiro compounds exhibit photochromism when irradiated by photons. These characteristics have been exploited to study liquid crystals. Shragina et al.^{[68](#page-53-0)} have studied the photochromic molecules (44) and (45), which

Galanthamine (43) , which contains the basic spiro $[5H-2-]$ benzazepine- 5^{\prime} , 1^{\prime}-cyclohexene] tricyclic fragment, is among some of the most widely used and most effective experimental drugs in the treatment of Alzheimer's disease, because of its potent acetylcholinesterase activities.⁵

3. Photochromism

Optical memory systems including molecular memory switches impact upon photochromism. $56-58$ It has been proposed^{[59,60](#page-53-0)} that information storage systems are based on a combination of two types of reversible process, namely photochemical cis/trans isomerisation and electrochemical The side-chain liquid-crystal polymers (46), (47) and (48) contain mesogenic and photochromic groups.[69,70](#page-53-0) The photochromic liquid-crystal polymers open up many possibilities in molecular engineering because the mesogenic and photochromic groups are more autonomous in this case.

Me

Me

Masafumi et $al.^{73}$ $al.^{73}$ $al.^{73}$ have reported that, on exposure of the spirobenzothiapyran 52 to UV light in polar solvents like methanol and acetone, the light-yellow solution turned to blue-green. The colour was spontaneously bleached at room temperature when the irradiation was turned off.

Photolysis of the spiro compound (53) results in photoisomerisation via the nitrile-stabilised biradical (54). Photolysis of the dienone (55) in the presence of tetramethylethylene gave the spiro compound $(57)^{74}$ $(57)^{74}$ $(57)^{74}$ through the biradical (56), as shown in [Scheme 2](#page-14-0).

4. Synthesis

It has been reported that the incorporation of a crown ether moiety into a spirobenzopyran affords ion-responsive photochromic materials.^{[71](#page-53-0)} The spirobenzopyrans (49) and (50) showed a selective binding ability to Mg^{2+} and Ag⁺ with negative and positive photochromism, respectively. Among the metal ions, only Ag^+ facilitated photoiso-merisation to the corresponding merocyanine form.^{[72](#page-53-0)} In the case of the spirobenzopyran 51, thermal isomerisation and facilitated photoisomerisation to the merocyanine form were observed in presence of Ag^+ .

A spiro compound contains two rings fused at a common point, mostly a carbon atom. The antithetic analysis of a spiro unit may result in a large number of possible prestructs, which can be used for the synthesis of target spiro molecules. The analysis can be undertaken by bond disconnection either at branch appendages (bond other than exocyclic) or ring appendages (exocyclic bond) or at both.⁷⁵ The process leads to the prediction of several synthon components, which include cyclic compounds on which a spiro unit can be generated or acyclic derivatives, which can cyclise to yield the target spiro compound.

56

Scheme 2.

4.1. Antithetic analysis of spiro unit

4.1.1. Cleavage at branch appendage(s). Bond disconnection at a bond of any ring of the spiro unit other than exocyclic will lead to a prestruct (\mathbf{Ia}) having a single synthon component with a ring and two alkyl units. In prestruct Ia, the cyclisation may occur due to the reconnection of the terminal point of the pendant alkyl groups to achieve the target.

55

 MeC

 MeC

Bond disconnections at two branch appendages in a single ring lead to the prestruct (Ib) with two synthon components, one being similar to Ia and the other an alkyl unit. Bond disconnection at branch appendages in both rings results in the generation of acyclic prestructs (Ic, Id and Ie). Prestruct Ic may have a single synthon component, whereas Id and Ie may have two and three acyclic components, respectively (Scheme 3).

Me 57

Scheme 3.

4.1.2. Cleavage at ring appendage(s). A spiro compound has four ring appendages having the possibilities of four types of bond disconnections (Scheme 4). A single bond disconnection in a single ring will lead to the formation of a prestruct (IIa) having a synthon component with a ring attached with a pendant alkyl group. Bond disconnection at two ring appendages of a single ring generates a prestruct (IIb) with two synthon components, one being a ring and the other an alkyl group. From this prestruct, the synthesis may occur either in a single step or in a tandem synthesis involving two steps. Disconnection at two ring appendages in two different rings may lead to an acyclic prestruct (IIc) , which contains all the atoms of the spiro-skeleton. The target molecule can be synthesised either in a single step or in a two-step process.

Three bond disconnections, two at one ring and one at the other ring, lead to the prestruct (IId) with two acyclic alkyl groups.

4.1.3. Disconnection at both branch and ring appendage(s). Bond disconnections at a ring appendage and a branch appendage in a single ring lead to the prestruct (IIIa) having synthon components, a ring with a pendant alkyl group and an acyclic alkyl fragment. Two bond disconnections in two different rings at one ring appendage and the Scheme 4. other at branch appendage will lead to a prestruct (IIIb) with

Scheme 6.

an acyclic synthon component having a branched alkyl group. Cleavage at two ring appendages and one branch appendage gives rise to the prestruct (IIIc) with all alkyl synthon components, while disconnections at two ring and two branch appendages engender a prestruct (IIId) with four synthon components [\(Scheme 5](#page-15-0)).

4.2. Synthesis of spiro compounds by using prestruct Ia

The prestruct Ia is due to bond disconnection at a branch appendage in a single ring. The corresponding synthon component has mostly a ring and two pendant alkyl groups attached at the carbon, which becomes the bridge head in the spiro nucleus. The synthesis of 63 from a spirogenic compound (60) is an example of the use of prestruct Ia. The synthon component 60, obtained from 58 through an intermediate 59, cyclises by an attack of a methylene unit

on a carbonyl group to yield 61, which, after derivatisation with ethylene glycol, cleaves to yield the spiro compound 62 and, on subsequent work up, produces (\pm) -acorenone B $(63)^{76}$ $(63)^{76}$ $(63)^{76}$ (Scheme 6).

Schobert and co-workers^{[77](#page-53-0)} have reported the synthesis of spirodiones 68 by using an α -hydroxy allyl ester (64) as the ring synthon component. Intramolecular Wittig olefination between 64 and the phosphorous ylide (65) afforded 66. Heating a solution of 66 in toluene to 180 \degree C in a sealed glass tube yielded 67, which undergoes autoxidation to the hemiketal endoperoxide (68) (Scheme 7).

A convenient synthesis of spirodiones 71 has been achieved by a one-pot tandem cyclisation–elimination process starting from the allyl-lacto acetals 70, prepared from the commercial ethyl 2-cyclohexanone carboxylates (69) [78](#page-53-0) (Scheme 8).

Scheme 7.

Scheme 9.

 $(-)$ -Sibirine (76) was synthesised from a chiral sulphoxide (72) used as a synthon component. The sulphoxide 72 was converted into 73 by the reaction of allylmagnesium bromide and subsequently into 74. The key transformation for this purpose was the conjugate addition Pummerer reaction^{79} involving the transformation of 74 into spiro compound 75 (Scheme 9).

When the chiral sulfoxide 77 was used instead of 72, the spiro compound 78 was formed as a single stereoisomer on reaction with allylmagnesium bromide. Compound 78 was converted into 79 on treatment with mercuric(II) trifluoroacetate, which underwent ring opening to form 80 in high yield. The spirodecanes 81 and 82 have been prepared from 80, as shown in Scheme 10.

Sharma et al. 80 have achieved the synthesis of spiro carbonlinked disaccharides (86) and (87) starting from a chiral ketone (83) and furan through 84 and 85, as shown in [Scheme 11](#page-18-0).

Ortho-substituted aminobenzenes (88) were used to synthesise spiro compounds containing a 2,3-dihydroindole unit (89), which occurs in the intermediates for the synthesis of Aspidosperma alkaloids, by using a steroselective 1,5 electrocyclisation, which can be brought about thermally or by deprotonation⁸¹ using t-BuOLi and t-BuOH [\(Scheme 12\)](#page-18-0).

Zubkov et al. 82 82 82 have synthesised the spiro compounds (94) and (95) by Wagner–Meerwein rearrangement of the diepoxy derivatives (93). Compound 93 was synthesised from furfurylamine (90) and cycloalkanones through 91 and 92, as shown in [Scheme 13.](#page-18-0)

Cossy et al.^{[83](#page-53-0)} have used a cyclic α -keto ester as the synthon component for the synthesis of spirodiols 98. The alkylation

Scheme 11.

 $NH₂$

90

and reduction of 96 generated the hydroxy esters 97, which, on cyclisation, furnished the spirodiols 98 [\(Scheme 14\)](#page-19-0).

4.3. Synthesis of spiro compounds by using prestruct Ib

The prestruct Ib is due to the bond disconnection at two branch appendages of one ring in the spiro unit. Therefore, the synthon component has one ring unit similar to prestruct Ia and the other an alkyl unit. A number of spiropyrans were Scheme 12. Scheme 12. Scheme 12. Synthesised by Padmavathi et al. 84 using prestruct Ib. They

95

 0^oC , 1h

 $\ddot{}$ H

'n

AcO AcO \overline{H}

 Ac

93

(i) PhH, Δ
(ii) allylMgBr/Et₂O

 H_2O_2 (50%) HCOOH Δ , 3h

AcO

 $n = 1, 2$

 $\mathbf H$

94

a
Ac 92

 $\lambda_{\rm n}$

 $\begin{matrix} 1 \\ 91 \end{matrix}$ $|Ac_2O, \Delta$

Scheme 14.

have taken the 2,6-diaryl-4,4-disubstituted-4H-pyrans (102) and (109) as the synthon components. The synthesis of 102 and 109 were achieved by the reaction of 99 with 100 and 107, respectively, through the formation of 101 and 108, respectively. Cyclocondensation of 102 and 109 with $NH₂NH₂·H₂O$, $NH₂OH·HCl$, $NH₂CONH₂$ and $NH₂CSNH₂$ in the presence of base afforded the desired spiro compounds 103–106 and 110–113, respectively, as shown in Schemes 15 and 16.

Prestruct Ib was also used by Coelho et al. 85 for the synthesis of spiro[thioxanthene-napththopyrans] (116). The spiro compounds were synthesised from substituted naphthols (115) and an intermediate, obtained from thioxanthone (114), using a one-step reaction [\(Scheme](#page-20-0) [17\)](#page-20-0). The photochromic properties of these compounds were studied under flash photolysis and continuous irradiation.

Kurth and co-workers^{[86](#page-54-0)} have reported the synthesis of spirofused (C_5) -isoxazolino- (C_4) -pyrazolones $(1-\alpha x^2,7,8-\alpha)$ -triazaspiro[4,4]-2,8-nonadien-6-ones) 119, by using β -ketoesters (117) as the ring synthon component. The reaction of 117 with hydrazine results in the intermediate hydrazone 118. This hydrazone undergoes cyclo-elimination to give the isoxazolinopyrazolones (119) [\(Scheme 18](#page-20-0)).

4.4. Synthesis of spiro compounds by using prestruct Ic

Bond disconnection at branch appendages of two different rings engenders prestruct Ic. The corresponding synthon component is an acyclic branched compound having a quaternary carbon. The synthesis of 4,9-dimethylspiro[4.4] nonane-2,7-dione (125) is an example of the use of prestruct Ic. The synthon component 120 was synthesised from acetylacetone. The 4-pentenal (121) was formed from 122 by monoacetylation and subsequent oxidation. The compound 123 undergoes Rh-mediated cyclisation twice to form the desired spirodiketone $(124)^{87}$ $(124)^{87}$ $(124)^{87}$ [\(Scheme 19\)](#page-20-0).

The tricyclic sila-alkane $(128)^{88}$ $(128)^{88}$ $(128)^{88}$ and bicyclic sila-alkane (129) were synthesised from bis(phenylethynyl)-silanes (126) through an intermediate (127), as shown in [Scheme 20](#page-21-0).

Scheme 18.

Scheme 17.

Scheme 16.

Scheme 20.

4.5. Synthesis of spiro compounds by using prestruct Id

The prestruct Id is due to bond disconnections at three branch appendages of two rings, that is, two bond disconnections at one ring and the third bond disconnection at the other ring. In this case, one of the synthon components is similar to prestruct Ic and the other component is an alkyl unit. To the best of our knowledge, no report is available describing the synthesis of spiro compounds by using synthon components corresponding to prestruct Id.

4.6. Synthesis of spiro compounds by using prestruct Ie

When the bond disconnections take place at four different branch appendages, the prestruct Ie is generated. An example of the use of prestruct Ie is the synthesis of spirobiscalix[4]crowns (132) from the synthon components 130 and 131, as shown in Scheme 21.^{[89](#page-54-0)}

Moll et al.^{[90](#page-54-0)} have also used pentaerythrityl tetrabromide (133) as one of the synthon components for the synthesis of spiro[3,3]heptane-2,6-dispirofluorene (134) (Scheme 22).

Scheme 22.

The novel spiro-compounds (139a–f) have been prepared by Schulte et al. 91 using pentaerythrityl tetrabromide as the synthon component. The synthesis has been carried in three steps by one-pot oxidative coupling of guaiacol (135) and 1,2-dialkoxybenzenes (136a–f), followed by dimethylation of triphenylenes (137) to 138 and subsequent etherification with tetrabromide [\(Scheme 23\)](#page-22-0).

4.7. Synthesis of spiro compounds by using prestruct IIa

The prestruct **IIa** is due to bond disconnection at a ring appendage in one ring. The corresponding synthon component has mostly a ring and an alkyl group attached at carbon, which subsequently becomes the spiro centre. Prestruct **IIa** is used in the synthesis of $(-)$ -perhydrohistrionicotoxin from 142. The synthon component 142, an amine, was synthesised from 140. Hydrolysis of the thioketal moiety of 140 by treatment with N-chlorosuccinimide (NCS) and silver nitrate gave 141, which was treated with triphenylphosphine to yield 142. Spirocyclisation of 142 was carried by treatment with dichlorotitanium(IV) diisopropoxide to furnish the spirocycle 143. Catalytic hydrogenation of the double bond in 143 resulted in perhydrohistrionicotoxin (144) ([Scheme 24\)](#page-22-0).⁹²

A number of tetraazaspiro[5,5]trione compounds (146) or (147) were synthesised from 145 in the presence of diethyl azodicarboxylate (DEAD) or NCS or N-bromosuccinimide

 $a: R = C_5H_{11}$; $b: R = C_6H_{13}$; $c: R = C_7H_{15}$; $d: R = C_8H_{17}$; $e: R = C_9H_{19}$; $f: R = C_{10}H_{21}$

Scheme 23.

(NBS) as dehydrogenating agents. The compounds 145 were synthesised from barbituric acid [\(Scheme 25](#page-23-0)).^{[93](#page-54-0)}

Kido et al. 94 have developed a new method for the synthesis of $(+)$ -acorenone B (151) through a spiro compound (150) based on spiroannulation of a cyclic arylsulphonium ylide (149) obtained from the diazo ketoester 148 ([Scheme 26](#page-23-0)).

Prestruct **IIa** has been used by Pigge et al., ^{[95](#page-54-0)} who have synthesised novel cyclohexadienyl azaspirocyclic ruthenium complexes (RuCP) (153) by using the corresponding N-benzyl acetoacetamide derivatives (152) as the synthon component. Demetalation of 153 to provide 154 has been accomplished by several oxidants, out of which $CuCl₂$ proved to be the most efficient ([Scheme 27\)](#page-23-0).

A synthon component corresponding to prestruct IIa has been used to form stabilised Meisenheimer-type salts^{[96](#page-54-0)} (156) by anionic spirocyclisation of 3-nitro-4-(2-hydroxyphenoxy)-2H-1-benzopyran-2-ones (155) ([Scheme 28\)](#page-23-0).

Spiro-oxindoles (159) and (160) have been synthesised from $\arcsin(158)$ by an asymmetric Heck reaction^{[97](#page-54-0)} using BINAP (157) . The synthon component

 $R = H$, 2'-Me, 3'-Me, 4'-Me, 2',3'-(Me)₂, Et, OMe, 2',3',4'-(OMe)₃; R_1 = Me; R_2 = H, Me; X = Cl, Br

Scheme 25.

Scheme 26.

158 undergoes palladium catalysed cyclisation to form the dispiro compounds (159) and (160), as shown in [Scheme 29](#page-24-0).

The formation of spiro [4.4] compounds 163 and 164 has been achieved by Mori et al.^{[98](#page-54-0)} by enantioselective spirocyclisation of the diene 161 in the presence of a catalyst 162 ([Scheme 30](#page-24-0)).

Pearson et al.^{[99](#page-54-0)} have explored the possibility of making thiol lactones (167) starting from an iron-complexed allyl thio ester (165) through corresponding spiro intermediate (166) as shown in [Scheme 31](#page-24-0).

In an effort to synthesise the biologically active spiro alkaloids, nitramine, isonitramine and sibirine, Cossy et al.^{[100](#page-54-0)} have chosen β -ketoamide (168) as a synthon component of the prestruct IIa for the preparation of chiral

 $R = H$, OMe

Scheme 27.

lactams (171) and (172). The ketoamide (168) was heated with 1 equiv of manganese(III) acetate in ethanol and a mixture of imines (169) and (170) was produced. These latter imines have been separated and hydrolysed with aqueous acetic acid to afford the spiro-lactams (171) and (172) ([Scheme 32](#page-24-0)).

A recent approach reported by $Simplins¹⁰¹$ $Simplins¹⁰¹$ $Simplins¹⁰¹$ involves the use of prestruct IIa to generate a range of spirocyclic ethers 174 of different ring size. Intramolecular conjugate addition of an alkyl radical derived from the phenylselenyl compound (173) to an enone system afforded the desired spiro ethers 174 ([Scheme 33\)](#page-24-0).

Scheme 29.

Scheme 33.

 \overline{O}

166

Martin-Lopej and Bermezo.¹⁰² have described the synthesis of the $1,6$ -diazaspiro[4.5]decane (178) and 6-aza-1oxaspiro[4.5]decane (179) systems by oxidative cyclisation of 176 and 177, respectively, which are obtained from an N-substituted tetrahydropyridine derivative (175) (Scheme 34).

Me₃NO, PhF

Scheme 30.

Scheme 31.

 $(CO)_{3}Fe$ $(CO)_{3}F$ hv, 350 nm, PhH

165

Naf et al.^{[103](#page-54-0)} have used an enone system **180** to prepare the spiro hydroxy ketone 181 in a single step. Conjugate addition of lithium dimethylcuprate to 180 and spontaneous intramolecular aldol condensation furnishes the desired spiro compound 181 with high yield ([Scheme 35](#page-25-0)).

 Ω

167

Scheme 34.

Scheme 35.

Similarly, Buchi et al.^{[104](#page-54-0)} have used a fulvene (182) to obtain a spiro compound 183 by using the same reagents, which, on subsequent treatment with N_2H_2 and H_2O_2 yields a single stereoisomer 184 in high yield (Scheme 36).

The synthesis of a spiro diketone (186) with good enantioselectivity (85%) has been achieved^{[105](#page-54-0)} from a cyclohexanone ring system 185 on treatment with a Lewis acid in the presence of optically pure (S, S) cyclohexane-1,2-diol by way of a Michael addition mechanism. The reaction when carried out in the presence of ethylene glycol forms the spiro compound 186 (55% of yield) along with a minor product 187 $(16\% \text{ of yield})^{106}$ $(16\% \text{ of yield})^{106}$ $(16\% \text{ of yield})^{106}$ (Scheme 37).

Iron-complexed allyl amides or esters (188 and 192) have been found to undergo thermally induced stereospecific spirocyclisation to afford spiro compounds (189 and 190, and 193 and 194, respectively), which on further demetalisation resulted in formation of (191) and (195), respectively, as shown in Scheme 38. [107](#page-54-0)

Scheme 36.

Scheme 37.

Scheme 38.

The products 190, 194 and the starting material undergo competing rearrangements of the complexed diene unit. When the optically pure amide 196 was taken as the starting material, these competing processes were suppressed by incorporating an electron-withdrawing CN group at C_5 to form 197, which yielded an enantiomerically pure spiro compound (198) [\(Scheme](#page-25-0) [39\)](#page-25-0).

Singh^{[108](#page-54-0)} has also used prestruct Π a in the oxidation of O -vanillyl alcohol (199) with aqueous sodium metaperiodate to produce a spiro intermediate (200) in the presence of cyclopentadiene in a biphasic (CH_2Cl_2/H_2O)

medium containing cetyltrimethyl ammonium bromide (CTAB) as a phase-transfer catalyst, furnished the adduct 201 in 70% yield (Scheme 40).

Yamamoto et al.^{[109](#page-54-0)} have synthesised α -acoradine (208) from a cyclohexane ring system 202 containing a long side chain by an intramolecular Sakurai–Hosomi reaction (Scheme 41). An allylsilane (204) has been made from 202 through an aldehyde intermediate (203) followed by cyclisation in the presence of ethylalumium dichloride to afford the spiro compounds (205) , (206) and (207) . Unfortunately, the stereoselectivity was poor and 207 is convertible into α -acoradine 208.

Scheme 40.

Scheme 41.

Scheme 43.

Scheme 44.

Ihara's group^{[110](#page-54-0)} has reported the synthesis of the sesquiterpenes (\pm)-erythrodiene (214) and (\pm)-spirojatamol (215) by the use of an intramolecular Mukaiyama aldol reaction, as shown in [Scheme 42.](#page-26-0) The acetal 209 when treated with $Me₃SiI/(Me₃SiI)₂NH$ gave a mixture of 210 and 211, which, on demethylation and oxidation, yielded the ketones 212 and 213. These latter ketones have been converted into 214 and 215.

Another method was used for the synthesis of the sesquiterpenes (\pm)-erythrodiene (214) and (\pm)-spirojatamol (215) by Huang and Forsyth.^{[111](#page-54-0)} The precursor

216 when allowed to react with a mercuric salt followed by photodimerisation gave a mixture of spiroketones out of which one derivative was elaborated into 214 and 215, as shown in Scheme 43.

Philip and Stephen^{[112](#page-54-0)} have reported the synthesis of a spirocyclic system (219) from the compound 217, which was converted to the spiro compound (218) through a free radical approach, as shown in Scheme 44.

Nishiyama et al.^{[113](#page-54-0)} have constructed the spirodienones (221) and (222) ; (224) ; (226) and (227) by aniodic oxidation

Scheme 46.

Scheme 47.

of the corresponding alcohol derivatives (220), (223) and (225), respectively, as shown in [Scheme 45](#page-27-0).

In the synthesis of manzamine-A, Brands and Dimichele^{[114](#page-54-0)} used 230 and 232 as the synthon components to synthesise a spirogenic compound (233) for the construction of the spirocentre in 234 with the desired stereochemistry. One of the synthon components 230 was prepared from 3-amino-1 propanol (228) through 229 and the other unit 232 was synthesised from a pyroglutamic acid derivative (231) (Scheme 46).

Tonghaosu (240a), a natural product isolated from several plants of the tribe Athemdeae, is a [4.4]spiroketal. The spiroketal was synthesised from furfuraldehyde (235) through a series of reactions involving intermediate $(236 - 239)$, as shown in Scheme 47.^{[115](#page-54-0)}

Craig et al.^{[116](#page-54-0)} have prepared 6.6- and 6.5-spiroketals (244–250) by acid-catalysed cyclisation of 242 and 243, which were obtained by alkylation and epoxide-opening reactions of 3,4-dihydro-6-[(p-toluenesulfonyl)methyl]-2Hpyran (241) [\(Scheme 48\)](#page-29-0).

Scheme 48.

Zhang et al.^{[117](#page-54-0)} have built a spiro centre in a cyclohexanone ring through a stepwise process using prestruct IIa. The precursor (253) of the spiro compound (255) was prepared by alkylating at the oxygen function of the cyclohexanone (252) using a substituted o -bromobenzoic acid (251). The cyclisation was proposed to proceed via a free-radical intermediate (254) , ^{[118](#page-54-0)} as shown in Scheme 49.

In the synthesis of spirotryprostatin, Edmonson and Danishefsky^{[119](#page-54-0)} have used a 6-methoxytryptophan (256) as one of the synthon components for the synthesis of the spiro-oxindole unit 260. The synthesis commenced with a Pictet–Spengler reaction^{[120](#page-54-0)} of the aldehyde 257 with 256 to yield 258, which on further hydroxybromination resulted in the spirogenic compound (259) ([Scheme 50](#page-30-0)). The spiro compound 260 serves as an intermediate for the synthesis of spirotryprostatin.

Semmelhack^{[121](#page-54-0)} has explored the synthesis of $(+)$ -acorenone and (\pm) -acorenone B [\(Scheme 51\)](#page-30-0) using the prestruct IIa. The diastereoisomers 262 and 263, on treatment of an anisole (261) with chromium hexacarbonyl, were converted into the spiroketones 264 and 265, respectively, through a variety of reactions, as shown in [Scheme 51](#page-30-0).

 (\pm) -Acorenone B (271) was synthesised stereoselectively by Trost et al.^{[122](#page-54-0)} by using a ketone 266 and cyclopropylidenediphenylsulphonium fluoroborate (267)

 $R_1 = H$, OMe, Boc; $R_2 = 2.4 - C1_2C_6H_3$, 2-ClC₆H₄, 2-CF₃C₆H₄, 4-MeOC₆H₄

Scheme 50.

Scheme 51.

through various intermediates (268–270), as shown in [Scheme 52](#page-30-0).

4.8. Synthesis of spiro compounds by using prestruct IIb

Prestruct **IIb** is due to bond disconnection at two ring appendages at one ring. The two synthon components consist of a ring and an alkyl group.

Adamantanone (272) and an amine 273 have been used as synthon components in the synthesis of spiro[piperidine- $2,2'$ -adamantane] $(276)^{123}$ $(276)^{123}$ $(276)^{123}$ as shown in Scheme 53. The reaction of 272 and 273 gave the expected product 274, which was converted into the dithiolane derivative (275). Subsequent hydrogenolysis of 275 furnished the desired spiro compound 276.

The same strategy was adopted for the synthesis of a $2,2'$ spiropiperidine skeleton (278) by taking cyclic ketones 277. Subsequent keto-deprotection resulted in the formation of 279 (Scheme 54).

Prestruct IIb has been used by Stork et al. 124 for the synthesis of (\pm) β -vetivone (285). One of the synthon components, enone 280, was alkylated with homoallylic dichloride (281), the other synthon. The spiroketone 284 was formed via 282 and 283. Addition of methyllithium to 283 and subsequent treatment with acid gave $(+)$ - β vetivone (285) (Scheme 55).

Posner et al. 125 have reported a similar method for the synthesis of (\pm) β -vetivone (285) starting from a lactone (286) instead of the enone 280. The lactone (286) on alkylation with the allylic-homoallylic dibromide (287) forms the spiro-lactone (288), which has been converted into (\pm) β -vetivone (285) by a series of reactions, as shown in [Scheme 56](#page-32-0). Asaoka et al.^{[126](#page-54-0)} used the same dihalide (287) to alkylate a different substrate 291 for the

Scheme 53.

 $n=2,3$

Scheme 54.

ОE1 OEt OEt i. LDA, HMPA, THI ii. LDA. $C1$ 280 281 $\frac{1}{C}$ 282 283 i. MeLi. Et.C ii. HCl **OEt** 285 284 (\pm) - β -vetivone

Scheme 56.

Scheme 57.

synthesis of (\pm) β -vetivone (285) through 292 and 293 (Scheme 57).

A spiro[4.5] skeleton 297 was constructed steroselectively by the reaction of a bis-Grignard reagent 295 and ketone 294 in the presence of $CuBr \cdot Me_2S$ through an intermediate 296 (Scheme 58). 127 127 127

A steroselective construction of spiro[4.5]decanones (301) was established by Koft and Smith¹²⁸ by spirocyclisation of 300, which was obtained by the reaction of enone 298 with Grignard reagent 299 (Scheme 59).

Burnell and co-workers^{[129,130](#page-54-0)} have synthesised a number of spiro[4.5]decanes, which have been employed as the key

Scheme 58.

Scheme 60.

Scheme 61.

Scheme 62.

step in the synthesis of various natural products. The construction of the simple spiro systems 306 and 307 can be achieved by using acetal 302 and a silylated acyloin (303) through a series of reaction furnishing intermediates 304 and 305 (Scheme 60).

The same group^{[131,132](#page-54-0)} has also reported the synthesis of spiro compound 311 by the reaction of a ketone 308 and a silyated compound 309 through the intermediate 310. A significant amount of the byproduct 312 was also formed (Scheme 61). This problem has been avoided^{[133](#page-54-0)} by using $BCl₃$ instead of $BF₃·Et₂O$ in a reaction of cyclohexanone (313) with 309, which traps the initial adduct as a cyclic ester 314 (Scheme 62). The spirocycle 315 was formed by treatment of the cyclic ester 314 with hydrogen fluoride in methanol, followed by TFA.

Some spiro-1,3-oxathiane derivatives (318) were obtained by the condensation reaction of cyclohexanones (316) with 3-mercapto-1-propanols $(317)^{134}$ $(317)^{134}$ $(317)^{134}$ (Scheme 63).

 $R = H$, Me; $R_1 = H$, Me, Ph, t-Bu

Scheme 63.

The investigation revealed that compound 318 ($R=Me$, R_1 = H) exhibits helical chirality (due to the spiro skeleton) and a virtual triligand chiral centre (belonging to the 1,3 oxathiane ring) whereas compounds 318 exhibit semiflexible structure.

Prestruct IIb has been used by Pardasani et al. 135 for the synthesis of spiro-oxazolidinone (321) and spiro-pyrrolidine derivatives (322–324). The reaction of the synthon components, isatin (319) and a secondary cyclic α -amino

Scheme 64.

acid, 1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid (320), gives rise to an azomethine ylide, which underwent $[3+2]$ cycloaddition reaction with various dipolarophiles affording an inseparable stereoisomeric mixture of spirooxazolidinone (321) and spiro-pyrrolidine derivatives (322–324), as shown in Scheme 64.

Oda et al.^{[136](#page-54-0)} have generated a spirocentre by using a cycloheptatriene derivative 325 and cycloheptanone. The rection of 325 and cycloheptanone yielded 326, which on Mukaiyama aldol reaction and Nazarov cyclisation furnished 327. Shapiro reaction of 327 yielded 328, which was converted to an azulenium ion 329 by hydride abstraction (Scheme 65).

Spirotetrahydropyridine derivatives (333) and (334) were synthesised diastereoselectively in good yield by the reaction of chalcones (332), azolin-5-ones (330) and (331), benzaldehyde and ammonium acetate in ethanol. When benzalacetone (335) was used, the reaction occurred only with isoxazolin-5-ones such as 330 leading to spirans $(336)^{137}$ $(336)^{137}$ $(336)^{137}$ ([Scheme 66](#page-35-0)).

Spiroketones (340) were constructed stereoselectively using cyclopentanone or cyclohexanone (337) and a substituted bromoepoxide (338) in the presence of potassium hydride ([Scheme 67\)](#page-35-0). The spiroketones (340) can be used as key intermediates for the synthesis of different alkaloids, viz $(-)$ -histrionicotoxin.¹³⁸

Cannone et al.^{[139](#page-54-0)} have used prestruct IIb in a stereoselective spirocyclisation of enones. Enone 341 was alkylated by the dibromide 342 to form the spiro

Scheme 66.

Scheme 67.

compounds 343 and 344. The spiro compound 343 can serve as a precursor to sesquiterpenes with a spiro[4.5] skeleton (Scheme 68).

The common intermediate (348), a spiro[4.5] system, can be utilised in the synthesis of α -vetispirene, β -vetispirene, b-vetivone and hinesole. The spiro compound (348) can be synthesised from the ketone $(345)^{140}$ $(345)^{140}$ $(345)^{140}$

through the intermediates 346 and 347, as shown in Scheme 69.

Nizamuddin et al.^{[141](#page-54-0)} have prepared several spiro-oxadiazolothiazoline and spiro-thiadiazolothiazoline derivatives (351) and (354) from cyclohexanone. 1-Aroyl-cyclohexanehydrazones (349) and 1-aryl-cyclohexane-thiosemicarbazones (352) when separately treated with mercaptoacetic

Scheme 68.

 $R = H$, 2-Me, 3-Me, 4-Me, 2MeO, 4-MeO, 4-C1

Scheme 70.

acid afforded 350 and 353, respectively. On subsequent treatment with concentrated H_2SO_4 spiro-oxadiazolothiazolines (351) and spiro-thiadazolothiazolines (354) were obtained (Scheme 70).

The synthesis and thermal rearrangement of spiro[2.4] hepta-1,4,6-triene (358) has been studied by Billups et al. 142 Photolysis of diazocyclopentadiene (355) in (2-bromovinyl)trimethylsilane (356) forms the intermediate 357, which, on treatment with CsF, gives the desired spiro compound 358 (Scheme 71).

Pujari et al. $143,144$ have synthesised spiro tetrazine compounds (360) by taking the 3-aryl/alkylindan-1-one (359) and thiocarbohydrazide as the synthon components (Scheme 72).

Joshi and co-workers have used prestruct IIb for the synthesis of several spiro compounds as given below.

Spiropiperidine (364) was obtained by the reduction of the 363, which was obtained by the reaction of 1-indanone (361) and an amine (362) (Scheme 73).^{[156](#page-55-0)}

The reaction of steroidal 6-ketones (365) with $R(-)$ -2aminobutanol in the presence of p-TsOH as catalyst affords selectively the respective steroidal $(6R)$ -spiro-4'-ethyl-1',3'-oxazolidines (366) (Scheme 74).^{[157](#page-55-0)}

Shamuzzaman et al.^{[158](#page-55-0)} have used prestruct IIa for the synthesis of cholest-5-en- $(3R)$ - Δ^{2-1} ',3',4'-oxadiazoline (369) by taking cholest-5-en-3-one (367) and semicarbazide hydrochloride as the synthon components. Compound 368 was obtained by the condensation of 367 with semicarbazide hydrochloride. The spiro compound 369 was synthesised by the cyclisation of 368 with acetic anhydride and pyridine [\(Scheme 75\)](#page-39-0).

Scheme 73.

4.9. Synthesis of spiro compounds by using prestruct IIc

The bond disconnections at the ring appendages of two different rings generate the prestruct IIc. The synthon component may be a long acyclic compound.

Nagashi et al.[159](#page-55-0) have used titanium-catalysed cascade carboalumination of trienes (370) and (374) to generate the spirobicycles (372), (373), (375) and (376), respectively, through an intermediate (371)[\(Scheme 76\)](#page-39-0).

Trost et al.^{[160](#page-55-0)} have reported the formation of two spiro centres in tandem by using prestruct IIc, which is an example of building two spiro units by tricyclisation. The synthon component 377 undergoes cyclisation in presence of palladium to form the dispiro compound 378 ([Scheme 77\)](#page-39-0).

 (E) - and (Z) -2-methoxycarbonylmethylene-1,6-dioxaspiro[4.5]decanes (381) and (382) have been synthesised from an equilibrium mixture of acyclic keto alcohol (379) and hemiacetal (380) via intermolecular conjugate addition^{[161](#page-55-0)} ([Scheme 78\)](#page-39-0).

The (E) isomer could be obtained in a 52:1 ratio under thermodynamically controlled conditions using t-BuOK in THF, whereas a catalytic amount of $Pd(OAc)$ resulted in the (Z) isomer in a 95:1 ratio.

Scheme 75.

Scheme 76.

Scheme 77.

4.10. Synthesis of spiro compounds by using prestruct IId

The prestruct IId is due to bond disconnections at three ring appendages of two rings, that is, two bond disconnections at one ring and the third bond disconnection at the other ring. In this case, the synthon components are two acyclic groups. No report has been available to us, however, describing the synthesis of spiro compounds by using synthon components corresponding to prestruct IId.

4.11. Synthesis of spiro compounds by using prestruct IIIa

The bond disconnections at a ring appendage and also a branch appendage at the same ring lead to prestruct IIIa.

A series of spiroheterocycles 384 were synthesised by the reaction of cycloalkylidenemalononitriles (383) with an active methylene compound, MeCOCH₂CO₂Et, as shown in Scheme 79.^{[162](#page-55-0)}

Scheme 79.

Scheme 80.

Some spiroheterocycles, 1-oxa-2,7-diazaspiro[4.4]non-2 ene-6,8-diones (386), were synthesised by the reaction of itaconimides (385) with nitrile oxides using prestruct IIIa (Scheme 80).^{[163](#page-55-0)}

An approach to the creation of a spirocentre in an asymmetric fashion by taking a prestruct IIIa has been reported by Moreto et al.[164](#page-55-0) This method involves nickelcatalysed addition of halomethylcycloalkanes (387) to one end of the triple bond and carbonylation of the other end of the acetylenic sulfoxide (388) to form 389 in a stereocontrolled manner. Demetallation of 389 gives a byproduct 390 and a spiro compound 391, which is converted to the desired spiro compound 392 (Scheme 81).

A similar study by the same workers has been reported^{[165](#page-55-0)} using an achiral acetylenic system (393) to synthesise spirocyclopentanones (394) by a similar type of intramolecular carbonylative cycloaddition (Scheme 82).

Scheme 82.

Functionalised spirocyclic tetrahydrofurans (399) have been obtained by Jones and Toutounji^{[166](#page-55-0)} by using prestruct **IIIa**. The synthon components, cycloalkanones 395 and ethyl acetoacetate (396), undergo an acetoacetate ester dianion– aldol reaction to form 397, which is transformed into an α -diazo- β -keto ester (398). The carbene O–H insertion of 398 engenders the spiro compound 399, as shown in [Scheme 83](#page-41-0).

Scheme 83.

 $R = COOEt$, H; $R_1 = Me$; $R_2 = H$, Me

Scheme 84.

Scheme 85.

Scheme 86.

The synthesis of spiro compounds 402 and 403 using prestruct IIIa has been reported by Gelmi et al.^{[167](#page-55-0)} The synthon components, 3-chloromethylenindolones (400) and dienes (401), undergo Diels–Alder cycloaddition reactions in the presence of an ethylaluminium dichloride catalyst to furnish the desired spiro compounds (Scheme 84).

Okada and co-workers^{[168](#page-55-0)} have reported the synthesis of the spiro adduct 406 as a single product by the reaction of dienophile (404) with the pure (E) isomer of piperylene (405) (Scheme 85).

Yadav and co-workers^{[169](#page-55-0)} have used prestruct **IIIa** for the synthesis of spirodiene (411). Baylis–Hillman reaction of 2-cyclohexenone (407) with formalin furnished 408, which underwent spirocyclocondensation with 2-mercaptoethanal to give the spiro-alcohol 409. Oxidation of 409 furnished an aldehyde 416, which on Wittig olefination furnished the desired spirodiene 411 (Scheme 86).

A convenient synthetic strategy for the synthesis of a new spiroheterocyclic system 417 has been developed by Campiani et al. 170 The target compound (417) has been prepared by using a tetrahydroisoquinoline derivative (412)

Scheme 88.

428

 $Ar =$

 \overline{C} Me σ Ar

430

 $R' = C_6H_5$, 4-Me C_6H_4 ,

Scheme 90.

and ethyl iodoacetate (413) as the starting materials through intermediates 414, 415 and 416 ([Scheme 87\)](#page-41-0).

In the synthesis of NK-1 receptor antagonists, Maligres et al.^{[171](#page-55-0)} have used prestruct \overline{II} for the synthesis of a spirobicyclic ether unit. Condensation of 2-phenylallyl bromide (418) with ketopiperidine (419) in the presence of zinc dust in THF furnished the homoallylic alcohol (420). Hydroboration/oxidation of 420 with BH_3 ·THF or $BH_3 \cdot Me_2S$ gave 421 or 422, respectively. Oxidation of 421 and 422 with PCC gave the spirobicyclic lactone units (423) and (425), respectively. Similarly reaction of 421 and 422 with NaHMDS afforded the spiro compounds (424) and (426) ([Scheme 88](#page-42-0)).

Keglevich and co-workers^{[172](#page-55-0)} have reported the synthesis of phosphorous-heterocycles (429) and (431) by a $[2+2]$ cycloaddition reaction of 2,3-dihydrophosphole oxides (427) or 1,2-dihydrophosphonil oxides (430) with dimethyl acetylenedicarboxylate (428) ([Scheme 89](#page-42-0)). In the spiro-1,2 oxaphosphetes (429) and (431) the phosphorous atom has a trigonal bipyramidal geometry. The 1,2-oxaphosphetanes are well-known intermediates for the Wittig reaction.^{[173](#page-55-0)}

Two spirocentres are generated by the 1,3-dipolar cycloaddition of 2,6-bis(aryl methylidene)cyclohexanones (432) and hydrazonyl chlorides (433). The reaction proceeded regioselectively affording the tetraazadispiro compounds (434) and $(435)^{174}$ $(435)^{174}$ $(435)^{174}$ (Scheme 90).

The spirodiones 439 and 440 were synthesised by Hayashi et al.^{[175](#page-55-0)} using 2-acetylcyclohexane (436) or 2-acetyltetralone (437) on reaction with allyl acetate in the presence of a chiral phosphine ligand (L^*) and subsequent cleavage of the double bond followed by Aldol condensation of the resulting aldehyde through an intermediate 438 (Scheme 91).

Carreira and co-workers^{[176](#page-55-0)} have synthesised spiro[pyrrolidin-3,3'-oxindoles] (443) and (444) by using spiro[cyclopropane-1,3'-oxindole] (441) as the ring synthon component. Treatment of 441 and N-alkylsulfonylaldimines (442) with a catalytic amount of MgI₂ in THF afforded the desired spiro compounds (443) and (444) (Scheme 92).

A number of diastereomeric spiro tricyclic nitroso acetals (446 and 447) have been synthesised by the reaction of

Scheme 91.

 R_2 = Et, Ph, 4-MeC₆H₄, 2-MeC₆H₄, 2-BrC₆H₄, 4-BrC₆H₄, 4-CF₃C₆H₄, 4-MeOC₆H₄, furyl

Scheme 93.

 $R = Me$, *n*-Bu, Bn, *t*-Bu, OMe

Scheme 94.

Scheme 95.

4-nitrosoxazoles (445) and ethyl vinyl ether in dichloromethane, 177 as shown in Scheme 93.

The syntheses of spiroheterocycles 450 and 451 have been achieved by Richard et al.^{[178](#page-55-0)} through the reaction of aldehydes 448 and the pyrone 449 by the use of Ac₂O and piperidine (Scheme 94).

Zimmer et al. 179 have reported that oxidation of 8-substituted xanthines (452) by m -CPBA to give the spiro compound 453, whereas the xanthines 454 underwent rearrangement under similar conditions to yield 455 (Scheme 95).

 $(+)$ -Pulegone (456) was used as the synthon component for the synthesis of spiro compounds 460 and 461. Pulegone has been alkylated with 457 to obtain the diastereomers 458

and 459, which, on thermolysis, underwent a Conia-type cyclisation to yield a separable mixture of 460 and 461 (Scheme 96). 180

Scheme 96.

Scheme 97.

Eilbracht et al.^{[181](#page-55-0)} have also used prestruct **IIIa** by using lactone (462) and allylmagnesium bromide (463) as the synthon components for the synthesis of spiroketal (465). The addition of 463 to lactone 462 leads to a hemiacetal 464 as a single diastereoisomer.^{[182](#page-55-0)} The intramolecular ringclosure reaction of 464 in the presence of Rh(CO)₂acac gives the spiroketal (465) (Scheme 97).

Scheme 98.

Knolker et al. 183 183 183 have used a similar prestruct to synthesise spirocyclopentanes (468) and (470) by the reaction of an allylsilane (467) with 2-alkylidenecycloalkanones (466) and disubstituted exomethylene compounds (469), respectively, as shown in Scheme 98.

Barluenga et al.^{[184](#page-55-0)} have used an α , β -unsaturated exocyclic chromium carbene complex (471) and butadienes (472) to construct spirolactones (473), (474) and (475) (Scheme 99).

Provencal and Leahy^{[185](#page-55-0)} have used an unsaturated aldehyde (476) for the synthesis of the spiro compound 479. Compound 476 has been subjected to copper-catalysed conjugate addition to 477, followed by treatment with Caro's acid to form the diester 478, which, on Dieckmann condensation, followed by decarboxylation, furnished 479 (Scheme 100).

During a study of spirobenzopyran^{[186](#page-55-0)} analogues as $5-HT_{1A}$ receptor ligands and potential anxiolytic agents, a series of spin (pyrrolidine and piperidine 2,3'(2'H)-benzopyrans] (483) were synthesised and evaluated for their serotonergic and dopaminergic activities. Michael condensation of benzopyrans 480 with methyl acrylate (481) in the presence

 $X = 1$ -morpholino, OSiMe₃

Scheme 99.

Scheme 101.

of benzyltrimethylammonium methoxide afforded the nitroester derivatives (482). Compounds 482 were reductively cyclised with Raney nickel in methanol to give oxospiropyrrolidine benzopyran derivatives (483) (Scheme 101).

Suemune et al.^{[187](#page-55-0)} have reported the synthesis of a series of optically active spirocyclic diones (488–490). When acetal (484) was alkylated with bromoesters (485), it gave rise to the generic diesters (486). Dieckmann condensation of 486, followed by decarboxylation, produced the spiroketones

(487). Acetal cleavage of 483 furnished the spirodiones (488–490) (Scheme 102).

Baker and co-workers^{[188](#page-55-0)} have explored one of the aborative synthetic routes for the synthesis of spiro compounds, which involved a Schmidt rearrangement. Schmidt rearrangement of 491 with sodium azide and trichloroacetic acid gave 493 and spiroisoindolyl cyclohexadienone (492) in 30% yield. The formation of 492 involves a self-immolative intramolecular chirality transfer from a biphenyl axis to a spirocentre (Scheme 103).

Scheme 102.

 $R = H$, Tosyloxy, 2-thienyl, C=CPh, C=CTMS

Scheme 104.

The syntheses of spiropyrrolidines (496) and (498–501) were achieved by Mazal et al.^{[189](#page-55-0)} from azomethine ylides (494) and (497). The 1,3-dipolar cycloaddition of 494 or 497 with α -methylene- γ -lactones (495) gave the desired spiro compounds 496 and 498–501, respectively, with various extents of stereoselectivity and regioselectivity, as shown in Scheme 104.

Pujari et al.^{[190](#page-55-0)} have also used prestruct **IIIa** for the synthesis of spiro compound $50²$ by the reaction of

Scheme 105.

3,4,5,6-tetrahydro-pyrimidine-2-thione (502) with dichloromalonate (503) (Scheme 105).

Joshi et.al.^{[191](#page-55-0)} have used prestruct **IIIa** for the synthesis of a spiro[azetidine-2,3'-3H-indole]-2',4(1H)-dione (506) by the reaction of anil (505) with ClCH₂COCl (Scheme 106).

Spiro $[3H-3,2]$ -thiazolidines] (509) were synthesised by the treatment of mercaptoacetic acid with indolylimines (508) under microwave irradiation. Imines (508) were prepared from indole-2,3-dione (507) and aromatic amines, as shown in Scheme 107. [192](#page-55-0)

Takshi and co-workers 193 have reported the synthesis of spiroheterocycles (512) by the cycloaddition of ϱ -chloranil (510) with tricarbonyliron complexes (511) ([Scheme 108\)](#page-48-0).

Sasaki et al. 194 have explored the synthesis of spiroadamantane–aziridine, –azetidine, –triazabicycloheptadione,

Scheme 106.

Scheme 108.

–isoxazoline and –pyrazoline derivatives by ionic or cycloaddition reactions of methyleneadamantane (513) or ethyl adamantylidenecyanoacrylate (515). When methyleneadamantane (513) is treated with PhCNO, it gives spiroisoxazoline (514), whereas treatment of ethyl adamantylidenecyanoacrylate with $CH₂N₂$ affords spiropyrazoline (516) (Scheme 109).

Scheme 109.

Dandia and co-workers 195 195 195 have reported the synthesis of spiro compound 520 and 521 using prestruct IIIa. Michael condensation of 3-dicyanomethylene-2H-indol-2-ones (517) with 2-thiohydantoin (518) or 4-hydroxy-2H-1benzopyran-2-one (519) affords spiro compounds 520 and 521, respectively, under microwave irradiation (Scheme 110).

The reaction of 3-benzoylcyanomethylidine-1(H)-indol-2one (522) with a number of active methylene compounds leads to the formation of spiro compounds 523–525, whereas the reaction of mercaptoacetic acid, an α -amino acid and hydrazine hydrate furnishes the spiro compounds 526 and 527, respectively. Compound 522 is generated by condensation of 1H-indole-2,3-dione with benzoyl acetato-nitrile ([Scheme 111\)](#page-49-0).¹⁹⁶

Arturo and co-workers 197 have studied the synthesis of thiaspirohexane (529) and azaspirohexane (530) by the cycloaddition of N-mesityl cyclopropylideneazomethine (528) with $Ph_2C = S$ and $PhN=C(CN)_2$, respectively. Compound 528 is synthesised by the reaction of mesitylcyclopropyl formimidoyl chloride with Me3COK in THF ([Scheme 112\)](#page-49-0).

Chen et al.^{[198](#page-55-0)} have reported an efficient synthesis of (\pm) alantrypinone (534) by a hetero Diels–Alder reaction of a novel pyrazine diene (531) with a 3-alkylideneoxindole (532) through a spiro intermediate (533) ([Scheme 113](#page-49-0)).

In a sequential process, an imine (535) prepared from isatin and n-butylamine undergoes cycloaddition with chloroacetyl chloride to afford a mixture of stereoisomeric

Scheme 111.

Scheme 112.

Scheme 113.

Scheme 114.

 α -chloro- β -lactams (536 and 537). A free radical reduction of the mixture with tris(trimethylsilyl)silane leads to the formation of a spirocyclic β -lactam (538) (Scheme 114).^{[199](#page-55-0)}

In the synthesis of $(-)$ -acorone and several spiroses-quiterpenes, Marx and Norman^{[200](#page-55-0)} have used enone 539 as one of the synthon components. Enone 539 underwent a Lewis acid-catalysed Diels–Alder reaction with isoprene to give 540–543 [\(Scheme 115\)](#page-50-0).

Schultz and Taveras^{[74](#page-53-0)} have reported the synthesis of spiro $[2,5]$ octa-1,4-dien-3-ones (549) from methyl

Scheme 115.

Scheme 116.

2-methoxybenzoate (544) and iodoacetonitrile, through different intermediates 545–548 as shown in Scheme 116.

4.12. Synthesis of spiro compounds by using prestruct IIIb

Bond disconnections at a ring appendage as well as a branch appendage at two different rings generate prestruct IIIb. The synthon component consists of a branched acyclic compound. Grigg et al.^{[201](#page-55-0)} have synthesised a series of $5/6$ and 5/12–17-membered bicyclospiro compounds using prestruct IIIb. The enynes (550) undergo regiospecific hydrostannylation to afford the α -vinylstannanes (551), which, on bis-cyclisation-anion capture, yield the spirocyles

(552) via 5-exo-trig cyclisation followed by sp^3 -sp² intramolecular Stille coupling (Scheme 117).

4.13. Synthesis of spiro compounds by using prestruct IIIc

When bond disconnections take place at two ring appendages of one ring and a branch appendage of another ring, prestruct IIIc is generated. The synthon components are all alkyl groups.

From open-chain 1,3-disubstituted acetone derivatives (553 and 557), Saul et al.^{[202](#page-55-0)} have synthesised spirobisoxazolidine (554–556) and spirobisimidazolidine derivatives (e.g., 558)

 $n = 1,2$; $X = O$, NCOPh, NSO₂Ph

Scheme 118.

Scheme 119.

by reaction with different reagents. From a prochiral 1,3-dichloroacetone (559), the chiral spiro derivative 560 and 561 were obtained as racemic mixtures, which were further resolved by using brucine (Scheme 118).

A similar strategy^{[203](#page-55-0)} has been followed for the synthesis of spirodilactones. A mixture of spirodilactones (563), (564) and (565) were synthesised using malonic acid (561) and an alkene (562) by heating at 70 \degree C in the presence of glacial acetic acid and manganese(III) acetate. The spiro centre is generated at the $-CH_{2}$ – of malonic acid (Scheme 119). Of the possible three stereoisomers 563, 564 and 565, the unsymmetrical species 564 was the major product.

4.14. Synthesis of spiro compounds by using prestruct IIId

The prestruct IIId is due to bond disconnections at two ring appendages and two branch appendages, that is, two bond disconnections at one ring and two bond disconnections at the other ring. In this case the synthon components are four acyclic groups. No report has been available to us, however, describing the synthesis of spiro compounds by using synthon components corresponding to prestruct IIId.

4.15. Synthesis of spiro compounds by using a different type of prestruct

The synthesis of a strained spirocycle $(567)^{204}$ $(567)^{204}$ $(567)^{204}$ via photoinduced SO_2 –N bond cleavage of 566 was carried out using a prestruct, which is due to three bond disconnections at three ring appendages of different rings. This is an unique example, which used a different type of prestruct (Scheme 120).

Scheme 121.

Some novel spirodiones (569) were synthesised using a different type of synthon component, 568. 1-Alkyl/aryl-3 amino-1H,3H-quinolin-2,4-diones (568) react with urea in boiling acetic acid to form the spiro compounds (569)^{[205](#page-55-0)} (Scheme 121).

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Stereoselective preparation of trifluoromethyl containing 1,4-oxathiolane derivatives through ring expansion reaction of 1,3-oxathiolanes

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Abstract—Trifluoromethyl containing 1,4-oxathiolanes are synthesized in excellent yields and high stereoselectivities from the expansion reactions of sulfur ylide intermediates, which were prepared from the reaction of 2-diazo-3,3,3-trifluoro-propionic acid methyl ester and 1,3-oxathiolanes in the presence of $Rh_2(OAc)_4$.

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1. Introduction

Considerable attention has been given to trifluoromethyl containing organic compounds as agrochemical and pharmaceutical agents due to their unique properties arising from altered electron density, acidity, and lipophilicity. $\frac{1}{1}$ Accordingly, the development of new methods for the synthesis of the trifluoromethyl containing organic compounds is continuous to be a an important area of research in agricultural, medicinal, and organic chemistry.² 1.4-Oxathiolanes are important heterocycles occurring in natural and medicinal molecules,^{[3](#page-61-0)} for example, RNA polymerase inhibitor tagetitoxin. 4 Although the preparation of nonfluoro-1,4-oxathiolane are well documented, however, the stereoselective ratio of the products mostly is still low, in particular to those containing quaternary carbons.^{[5](#page-61-0)} Furthermore, it is difficult to introduce fluorine atoms into 1,4-oxathiolanes.^{[6](#page-61-0)}

2. Results and discussion

As part of a project on synthesis of fluorine-containing molecules using fluorinated diazocompounds, we wish to develop a method to synthesize trifluoromethyl containing 1,4-oxathiolane through ring expansion of 1,3-oxathiolane started from 2-diazo-3,3,3-trifluoro-propionic acid methyl ester.

Porter et al. reported 1,4-oxathiolanes could be synthesized from the reaction of 1,3-oxathiolanes and a silylated diazoacetate in the presence of a copper catalyst in moderate yield and low diastereomeric ratio. 5 In this paper, we will report the stereoselective synthesis trifluoromethyl containing 1,4 oxathiolane from the reaction of 2-diazo-3,3,3-trifluoropropionic acid methyl ester and 1,3-oxathiolanes (Scheme 1).

1,3-Oxathiolane 2 can be readily obtained from the reaction of 2-mercapto ethanol and carbonyl compounds in the presence of CAN in excellent yields. Initial studies were focus on the reaction of 1,3-oxathiolanes 2 derived from aromatic aldehydes and diazocompounds 1. To a refluxing benzene solution of 1,3-oxathilane 2 and catalyst was added a benzene solution of diazocompound 1 in 2 h. The stirring continues for an additional 2 h. $Rh_2(OAc)_4$ is a superior catalyst than the copper catalysts, for example, $Cu (acac)_2$, which is contradict to the results reported by Porter.^{[5](#page-61-0)}

Keywords: Stereoselective synthesis; Diastereomeric ratio; Lipophilicity; Sulfur ylide; Diazo; Trifluoromethyl; 1,4-Oxathiolane.

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The reaction gives 1,4-oxathiolanes 3 in high yields with excellent diastereomeric ratio. 1,3-Oxathiolanes derived from the aromatic aldehydes substituted with electronwithdrawing groups lead to the products in relatively higher yields than those electron-donating ones, but with lower diastereometric ratio (Table 1, entries 6–8). Chemical shift data from the NMR spectrum of the products suggests that the major isomer is the one in which the carbmethoxy and phenyl groups are trans-oriented. Those electron-rich substrates give 1,4-oxathiolanes in higher trans/cis ratio than those electron-poor ones. For example, it gives to almost exclusively trans diastereoselective products when the aromatic ring was substituted with methoxy group (Table 1, entry 8). The nitro-substituted substrate, however, it gives only 3:2 trans/cis diastereoselective ratio (Table 1, entry 8).

Table 1.

^a All products were fully characterized by spectroscopic methods. The yields were isolated yields.

 b Determined by $1H$ NMR integration of the reaction mixture.

The relative stereochemistry of 3 was further confirmed by a X-ray crystal diffraction analysis of compound 3a (Scheme 2). The structural analysis clearly shows the 1,4 oxathiolane ring of 3a keeps in a thermodynamic stable chair conformation. The large groups, methoxyl carbonyl and phenyl ring, lie in the equatorial position of six-member ring. Trifluoromethyl group, smaller than methoxyl carbonyl group, situates in the axial position. What resulted in the low diastereomeric ratio (2:1) with regard to the reaction of 2-(4-nitro-phenyl)-1,3-oxathiolane with 1,4-oxathiolane? We know that the trifluoromethyl and aromatic ring are in cis position according to the crystallographic study. The electrostatic repulsion between the trifluoromethyl and nitrophenyl, an electron-deficient group, makes the relatively bulky methoxyl carbonyl group tend to be axial set, in order to reduce such repulsion.

Scheme 2. The molecular structure of 3a.

Spiro structures existed in many medicinally or biologically important molecules.[7](#page-61-0) How to efficiently synthesize such compounds is still a challenge subject. It is in particular difficult to introduce fluorine atom into spiro-molecules. Spiro 1,3-oxathiolanes can be easily available from cyclic ketone with 2-mercapto ethanol. Based on the above experiments, the [1,2]-rearrangement of the intermediate sulfur ylides of spiro 1,3-oxathiolanes should give the trifluoromethyl containing spiro 1,4-oxatiolane. Under the same reaction conditions, the spiro compound 6 was indeed obtained in 59% yield, which was in line with our initial speculations (Scheme 3).

Scheme 3.

For comparison, 1,3-oxathiolane 5 was treated with trifluoromethyl diazoacetate 1 under similar reaction conditions, no [1,2]-rearrangement of intermediate sulfur ylides was not observed. The elimination product 7 was dominant (48%) (Scheme 4). The reaction might be preceded through a five-membered ring intermediate.

In summary, we have developed an efficient method for the stereoselective synthesis of trifluoromenthyl 1,4-oxathiolane through ring expansion of 1,3-oxathiolane ylide using

trifluoromethyl diazoactetate in the presence of $Rh_2(OAc)_4$. This catalytic, stereoselective and mild reaction will be the method of choice in many instances.

3. Experimental

Melting points were measured on a Temp-Melt apparatus and are uncorrected. Solvents were dried before use. ${}^{1}H$, 19 F, and 13 C NMR spectra were recorded on a Varian-360L instrument or Bruker DRX-400 spectrometer with TMS and TFA (δ CFCl₃= δ TFA+76.8) as the internal and external standards and the upfield as negative. IR spectra were obtained with an IR-440 Shimadzu spectrophotometer. Low-resolution mass spectra and high-resolution mass spectra (HRMS) were obtained on a Finnigan GC–MS 4021 and Finnigan MAT-8430 instrument, respectively. The X-ray structural analysis was performed with a Rigaku/AFC 7R Diffractometer. Elemental analyses were performed by this Institute.

3.1. General procedure for ring expansion

A mixture of 1,3-oxathiolane (2) (1 mmol) and $Rh_2(OAc)_4$ (5 mg, 0.01 mmol) in dry benzene (2 mL) was heated to reflux under a nitrogen atmosphere. A solution of 2-diazo-3,3,3-trifluoro-propionic acid methyl ester (1) (201 mg, 1.2 mmol) in benzene (2 mL) was added dropwise over 2 h through syringe. Reflux was continued for 2 h, and then the mixture was allowed to cool to rt. The solvent was removed under reduced pressure and the residue was purified by a flash chromatography on silica gel (petroleum–ethyl acetate) to give the ring expansion product 1,4-oxathiolane (3). The procedure for the preparation of products 6 and 7 are similar to that of 3.

3.1.1. 2-Phenyl-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3a). Colorless crystal with mp: 59–61 °C, yield: 93%, IR (KBr): 2987, 2957, 2924, 1747, 1498, 1455, 1432, 1253, 1164 cm⁻¹. ¹H NMR (300 MHz, CDCl₃): δ 2.44 (d, 1H, J=13.5 Hz), 3.41 (t, 1H, $J=12.3$ Hz), 3.80 (s, 3H), 4.07 (t, 1H, $J=11.7$ Hz), 4.49 (d, 1H, J=11.4 Hz), 5.49 (s, 1H), 7.32–7.46 (m, 5H) ppm. ¹³C NMR (75 MHz, CDCl₃): δ 24.7, 53.4, 55.8 (d, J_{F–C}= 24.8 Hz), 69.5, 82.4, 125.1 (q, $J_{F-C} = 283.3$ Hz), 127.2, 128.0, 128.5, 137.5, 166.1 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -59.89 (s) ppm. EI-MS (*mlz*, %): 306 (M⁺, 6), 200 (100), 172 (21), 105 (42), 77 (15), 59 (21). Anal. Calcd for $C_{13}H_{13}F_3O_3S$: C; 50.98, H; 4.28%. Found: C; 51.10, H; 4.25%.

X-ray data of 3a

 $C_{13}H_{13}F_3O_3S$: $M_w = 306.29$, CCDC no. 284941, orthorhombic, space group: $P2(1)2(1)2(1)$, $a=6.755(10)$ Å $b=$ 7.867(11) A, $c = 12.937(18)$ A; $\alpha = 91.464(3)^\circ$, $\beta = 92.651(3)^\circ$, $\gamma = 108.662(2)$ °; V=650.06(16) Å³, Z=1, D_c=1.794 g/cm³, $F(000) = 348$. Radiation, Mo K α ($\lambda = 0.71073$ Å). Crystal dimension, $0.58 \times 0.44 \times 0.34$ mm.

Intensity data were collected at 293(2) K with a Bruker P4 four-circle diffractometer with graphite monochromator and Mo K α radiation (λ =0.71073 A). A total of 8147

independent reflection were measured in range $2.36 < \theta <$ 27.0° . The structure was solved by directed methods and expanded using Fourier techniques. The nonhydrogen atoms were refined anisotropically, hydrogen atoms were included but not refined. The final cycle of fullmatrix least-square refinement was base on F^2 . The final R and wR value were 0.0490 and 0.1128, respectively. All calculations were performed using the SHELX-97 program.

3.1.2. 2-(2-Fluoro-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3b). Colorless crystal with mp: 71-73 °C, yield: 65%, IR (KBr): 3001, 2955, 2923, 1750, 1491, 1458, 1434, 1258, 1164 cm⁻¹. ¹H NMR (300 MHz, CDCl₃): δ 2.48 (d, 1H, J = 13.2 Hz), 3.43 $(t, 1H, J=15.0 \text{ Hz})$, 3.77 (s, 3H), 4.06 (t, 1H, $J=12.0 \text{ Hz}$), 4.49 (d, 1H, $J=11.7$ Hz), 5.48 (s, 1H), 6.94–7.59 (m, 4H) ppm. ¹³C NMR (75 MHz, CDCl₃): δ 25.1, 53.6, 57.2 (d, $J_{\text{F-C}}$ = 26.5 Hz), 70.0, 82.3, 114.4, 114.7, 123.9, 129.1 (q, $J_{\text{F-C}}$ = 283.4 Hz), 129.5, 130.8, 130.9, 165.0 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -60.3 (s, 3F) -116.8 (s, 1F) ppm. EI-MS (m/z , %): 324 (M⁺, 6), 200 (100), 172 (23), 123 (46), 113 (12), 59 (45). Anal. Calcd for $C_{13}H_{12}F_4O_3S$: C; 48.15, H; 3.73%. Found: C; 47.97, H; 3.84%.

3.1.3. 2-(2-Chloro-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3c). Colorless crystal, mp: 69-71 °C, yield: 75%, IR (KBr): 3011, 2958, 2967, 1934, 1740, 1478, 1437, 1258, 1160 cm⁻¹. ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$: δ 2.49 (d, 1H, J = 12.6 Hz), 3.47 (t, 1H, $J=13.5$ Hz), 3.75 (s, 3H), 4.06 (t, 1H, $J=11.7$ Hz), 4.49 (d, 1H, $J=11.7$ Hz), 5.59 (s, 1H), 7.25–7.65 (m, 4H) ppm. ¹³C NMR (75 MHz, CDCl₃): δ 25.7, 54.0, 58.1, 70.2, 79.8, 125.4 (q, $J_{\text{F-C}}$ =282.9 Hz), 126.4, 128.5, 129.8, 130.5, 132.8, 135.3, 164.7 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -59.7 (s, 3F) ppm. EI-MS (m/z, %): 340 (M⁺, 2), 200 (100), 172 (20), 139 (27), 113 (11), 59 (34). Anal. Calcd for $C_{13}H_{12}CIF_3O_3S$: C; 45.82, H; 3.55%. Found: C; 45.85, H; 3.62%.

3.1.4. 2-(2-Bromo-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3d). Colorless crystal, mp: 97–99 °C, yield: 75%, IR (KBr): 3062, 2995, 2920, 1745, 1474, 1431, 1262, 1225, 1170, 1153, 1095 cm⁻¹.
¹H NMP (300 MHz, CDCL): $\frac{5}{2}$ 2.8 (d, 1H, 1–13.8 Hz) ¹H NMR (300 MHz, CDCl₃): δ 2.48 (d, 1H, J = 13.8 Hz), 3.46 (t, 1H, $J=12.0$ Hz), 3.75 (s, 3H), 4.05 (t, 1H, $J=$ 12.0 Hz), 4.49 (d, 1H, $J=11.7$ Hz), 5.59 (s, 1H), 7.15–7.56 (m, 4H) ppm. 13 C NMR (75 MHz, CDCl₃): δ 25.7, 54.1, 58.0, 70.2, 82.4, 123.6, 125.5 (q, $J_{F-C} = 285.1$ Hz), 126.9, 130.1, 130.9, 131.8, 136.9, 164.6 ppm. 19F NMR (282 MHz, CDCl₃): δ -59.5 (s) ppm. EI-MS (*mlz*, %): 340/342 (M⁺ 1/1), 200 (100), 185 (15), 183 (16), 172 (19), 113 (8), 59 (29). Anal. Calcd for C₁₃H₁₂BrF₃O₃S: C; 40.54, H; 3.14%. Found: C; 40.69, H; 3.27%.

3.1.5. 2-(4-Bromo-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3e). Colorless crystal, mp: 62–64 8C, yield: 92%, IR (KBr): 2996, 2957, $1749, 1489, 1338, 1249, 1173, 1162, 1099$ cm⁻¹. ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$: δ 2.44 (d, 1H, J = 13.5 Hz), 3.37 (t, 1H, $J=12.3$ Hz), 3.78 (s, 3H), 4.02 (t, 1H, $J=11.7$ Hz), 4.46 (d, 1H, $J=11.7$ Hz), 5.41 (s, 1H), 7.24–7.26 (m, 2H), 7.41–7.44 (m, 2H) ppm. ¹³C NMR (75 MHz, CDCl₃): δ 21.0, 53.7, 55.3, 69.4, 85.8, 125.0 (q, $J_{F-C} = 283.4$ Hz),

129.1, 131.0, 131.4, 136.6, 166.0 ppm. 19F NMR (282 MHz, CDCl₃): δ -59.8 (s) ppm. EI-MS (*m/z*, %): 340/342 (M⁺, 4/4), 200 (100), 185 (26), 183 (24), 172 (21), 113 (17), 77 (7), 59 (35). HR-EI-MS calcd for $C_{13}H_{12}BrF_3O_3S$ (M⁺): 383.9643; found: 383.9657.

3.1.6. 2-(4-Nitro-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3f). Colorless crystal, mp: 77-79 °C, yield: 100%, IR (KBr): 3000, 2961, 2914, 1751, 1744, 1608, 1517, 1350, 1251, 1157, 1104, 1032 cm^{-1} . ¹H NMR (300 MHz, CDCl₃): δ 2.50 (d, 1H, J = 13.8 Hz), 3.42 (t, 1H, $J=12.3$ Hz), 3.83 (s, 3H), 4.06 (t, 1H, $J=12.0$ Hz), 4.52 (d, 1H, $J=11.7$ Hz), 5.57 (s, 1H), 7.58–7.61 (m, 2H), 8.14–8.18 (m, 2H) ppm. 13C NMR (75 MHz, CDCl3): d 24.7, 53.8, 55.2, 69.3, 80.9, 123.0, 126.8, 130.5, 144.6, 147.8, 165.9 ppm. 19F NMR (282 MHz, CDCl₃): δ -59.7 (s) ppm. EI-MS (*mlz*, %): 351 (M⁺, 1), 200 (100), 172 (17), 113 (3), 77 (1), 59 (5). Anal. Calcd for $C_{13}H_{12}NF_3O_5S$: C; 44.45, H; 3.44, N; 3.99%. Found: C; 44.36, H; 3.27, N; 3.89%.

3.1.7. 2-(4-Methyl-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3g). Pale yellow liquid, yield: 88%, IR (KBr): 3009, 2958, 2865, 2253, 1749, 1715, 1616, 1516, 1437, 1363, 1264, 1224, 1171, 1105 cm⁻¹.
¹H NMP (300 MHz, CDCL): $\frac{\delta}{2}$ 2.35 (c, 3H) 2.44 (d, 1H) ¹H NMR (300 MHz, CDCl₃): δ 2.35 (s, 3H), 2.44 (d, 1H, $J=12.6$ Hz), 3.42 (t, 1H, $J=11.2$ Hz), 3.80 (s, 3H), 4.06 $(t, 1H, J=11.7 \text{ Hz})$, 4.49 (d, 1H, $J=12.0 \text{ Hz}$), 5.45 (s, 1H), 7.12–7.15 (m, 2H), 7.26–7.35 (m, 2H) ppm. 13C NMR (75 MHz, CDCl3): d 21.2, 24.7, 53.5, 55.8, 69.6, 82.4, 123.3, 127.1, 128.6, 134.6, 138.2, 166.2 ppm. 19F NMR (282 MHz, CDCl₃): δ -59.9 (s) ppm. EI-MS (*m*/z, %): 320 $(M⁺, 8)$, 200 (100), 172 (15), 119 (50), 91 (17), 84 (19), 59 (27). HR-MALDI-MS calcd for $C_{14}H_{15}F_3O_3SNa$ $(M+Na^+)$: 343.0592; found: 343.0593.

3.1.8. 2-(4-Methoxyl-phenyl)-3-trifluoromethyl-1,4-oxathiinane-3-carboxylic acid methyl ester (3h). Pale yellow solid with mp: 82–84 °C, yield: 83%, IR (KBr): 3021, 2982, 2841, 1743, 1616, 1515, 1436, 1293, 1261, 1171, 1156, 1103 cm^{-1} . ¹H NMR (300 MHz, CDCl₃): δ 2.45 (d, 1H, J= 13.5 Hz), 3.41 (t, 1H, $J=12.0$ Hz), 3.80 (s, 6H), 4.06 (t, 1H, $J=11.7$ Hz), 4.49 (d, 1H, $J=13.5$ Hz), 5.41 (s, 1H), 6.82–6.85 (m, 2H), 7.27–7.31 (m, 2H) ppm. 13C NMR (75 MHz, CDCl3): d 24.8, 53.4, 55.2, 55.7, 69.6, 82.1, 113.2, 125.2 (q, $J_{\text{F-C}} = 281.2 \text{ Hz}$), 128.5, 129.6, 159.6, 166.2 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -59.8 (s) ppm. EI-MS $(m/z, %)$: 336 $(M⁺, 18)$, 200 (100), 172 (16), 136 (98), 135 (82), 113 (10), 77 (9), 59 (21). Anal. Calcd for $C_{14}H_{15}F_3O_4S$: C; 50.00, H; 4.50%. Found: C; 50.00, H; 4.47%.

3.1.9. 5-Trifluoromethyl-1-oxa-4-thia-spiro[5.5]undecane-5-carboxylic acid methyl ester (6). Pale yellow liquid, yield: 59%, IR (KBr): 3403, 2959, 2881, 1750, 1708, 1440, 1354, 1282, 1252, 1147, 1111 cm⁻¹. ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$: δ 1.38–1.87 (m, 6H), 2.01–2.05 (m, 2H) 2.30–2.34 (m, 2H), 2.86–3.04 (m, 2H), 3.76 (s, 3H), 3.86 (t, 1H, $J=5.7$ Hz), 4.18 (t, 1H, $J=8.1$ Hz) ppm. ¹³C NMR (75 MHz, CDCl₃): δ 22.6, 22.6, 27.6, 32.9, 53.2, 60.0, 77.1, 124.0 (q, $J_{\text{F-C}}$ =277.7 Hz), 165.7 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -52.1 (dd, $J^1 = 10.1$ Hz, $J^2 = 36.7$ Hz) ppm. EI-MS (*mlz*, %): $298(M^+,3),200(100),188(20),159(18),142(71),91(29),59$ (53) , 45(64). HR-MALDI-MS calcd for $C_{12}H_{17}F_3O_3SNa(M+$ Na⁺): 321.0748; found: 321.0759.

3.1.10. 3,3,3-Trifluoro-2-[2-(1-phenyl-vinyloxy)-ethylsulfanyl]-propionic acid methyl ester (7). Pale yellow oil (152 mg, 48%), IR (KBr): 3467, 2959, 1751, 1713, 1686, 1439, 1361, 1269, 1147, 1111 cm⁻¹. ¹H NMR (300 MHz, CDCl₃): δ 3.21 (q, 2H, J=4.8 Hz), 3.80 (s, 3H), 3.82 (s, 1H), 4.10 (t, 2H, $J=6.0$ Hz), 4.23 (d, 1H, $J=3.0$ Hz), 4.72 (d, 1H, $J=3.3$ Hz), 7.34–7.62 (m, 5H) ppm. ¹³C NMR $(75 \text{ MHz}, \text{CDCl}_3)$: δ 31.4, 49.7, 53.2, 63.2, 83.2, 127.8 (q, $J_{\text{F-C}}$ = 282.8 Hz), 125.0, 125.4, 128.2, 128.6, 159.6, 165.6 ppm. ¹⁹F NMR (282 MHz, CDCl₃): δ -52.5 ppm. EI-MS $(m/z, \%): 321 (M^+, 15), 201 (100), 179 (6), 141 (17),$ 77 (5), 59 (8). HR-MALDI-MS calcd for $C_{14}H_{15}F_3O_3SNa$: 343.0592; found: 343.0598.

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Callipeltins F–I: new antifungal peptides from the marine sponge Latrunculia sp.

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Abstract—Four new antifungal peptides, callipeltins F–I, were isolated from the marine sponge *Latrunculia* sp., collected off Vanuatu islands. Their structures were elucidated by NMR and MS analysis. The new callipeltins exhibited anti Candida activity in the 10^{-4} M range. Q 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Callipeltin A (1) and its congeners were isolated in our laboratories from the sponges Callipelta sp.^{[1,2](#page-69-0)} and Latrunculia sp. 3 Callipeltin A, that contains some unusual structural features such as the presence of numerous nonribosomial amino acids and a unique N-terminal aliphatic hydroxy acid moiety, represents the first member of a class of potent antiviral marine peptides. This growing family comprises papuamides A–D, obtained from Papua New Guinea collections of the sponge Theonella

mirabilis and *T. swinhoei*,^{[4](#page-69-0)} microspinamide from the Indonesian sponge Sidonops microspinosa^{[5](#page-69-0)} and neamphamide, from the Papua New Guinea sponge Neamphius huxleyi,^{[6](#page-69-0)} all sharing more or less a degree of structural homology suggesting a common pharmacophore. Recently, callipeltin A (1) was found to be a potent inotropic agent making it of interest as a regulator of myocardial contractility.[7,8](#page-69-0) The combination of interesting biological activity, unsual aminoacids and complex molecular architecture have attracted the interest of the synthetic chemistry community. ⁹⁻¹⁹

Keywords: Callipeltin; Peptides; Antifungal.

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These synthetic efforts have enabled the confirmation of the absolute stereochemistry of the 3,4-dimethylglutamine residue,^{[10](#page-69-0)} the revision of the 3-hydroxy-2,4,6-trimethylheptanoic acid end group, $16-18$ and the definition of the absolute stereochemistry of the β -OMeTyr unit.^{[19](#page-69-0)}

Callipeltin C (2), which is the acyclic derivative of callipeltin A, displays antifungal activity against Fusarium oxysporum, Helminthosporium sativum, Phytophtora hevea, and Candida albicans.

The absolute configurations of D-Ala, D-Arg and two D-alloThr residues were determined by LC–MS analysis of the acid hydrolysate derivatized with Marfey's reagent $(1-fluoro-2.4-dinitrophenvl)-5-L-alaninamide; L-FDAA)²⁰$ $(1-fluoro-2.4-dinitrophenvl)-5-L-alaninamide; L-FDAA)²⁰$ $(1-fluoro-2.4-dinitrophenvl)-5-L-alaninamide; L-FDAA)²⁰$ and comparison with appropriate amino acid standards. To establish the absolute configuration of 3,4-diMeGln and AGDHE residues, an authentic sample of callipeltin A was hydrolysed and derivatised with L-FDAA. The configuration of the remaining stereocentres in callipeltin F was assumed to be the same of callipeltin A.

Callipeltin C (**2**)

In order to re-isolate further amounts of callipeltins for additional pharmacological studies, we re-examined the polar extracts of the sponge Latrunculia sp., still available in our laboratories in large amounts. After a careful sequential HPLC separation on Vydac and Thermo HyPurity columns, we were able to isolate minor callipeltin-related open-chain derivatives, named callipeltins F–I (3–6).

In this paper, we describe the isolation and the structure determination of the new compounds.

2. Results and discussion

The lyophilised sponge was extracted with methanol and the crude methanolic extract was subjected to a modified Kupchan's partitioning procedure. Fractionation of the butanol-soluble material (ca. 4 g) by DCCC (CHCl $_3$ / $MeOH/H₂O$, 7:13:8 ascending mode) followed by repeated reversed phase HPLC afforded pure callipeltins F–I (3–6).

Callipeltin F (3) was obtained as colourless amorphous solid and its molecular formula was determined to be $C_{42}H_{79}N_{13}O_{14}$ by HR ESIMS. Extensive analysis of the ¹H and ¹³C NMR data of 3, including ¹H-¹H COSY, HMQC, HMBC spectra (see [Table 1](#page-64-0)), by comparison with those of callipeltin $C(2)$, disclosed the presence of one residue each of alanine (Ala), arginine (Arg), 3,4 dimethylglutamine (diMeGln), 4-amino-7-guanidino-2,3 dihydroxyheptanoic acid (AGDHE), 3-hydroxy-2,4,6-trimethylheptanoic acid, and of two residues of threonine (Thr). The amino acid sequence of 3 and placement of acyl substituent were assigned from the analysis of the fragmentation pattern in the ESI MS/MS mass spectrum [\(Fig. 1\)](#page-65-0).

The molecular formula $C_{54}H_{100}N_{16}O_{17}$ of callipeltin G (4) was deduced by HR ESIMS $[m/z \ 1245.7532 \ (M+H)^+]$. COSY, HOHAHA, HMQC, HMBC experiments readily disclosed the presence, in addition to the same residues found in callipeltin F of one residue of leucine (Leu) and N-methylglutamine (MeGln). The sequencing of these units by the analysis of ESI MS/MS fragmentation peaks led to the structure as shown in [Figure 2](#page-65-0).

The stereochemistry of the amino acid residues was determined by Marfey's method to be D-alloThr, D-Arg, L-Leu, L-MeGln, D-Ala.

Callipeltin H (5) analyzed for the molecular formula $C_{68}H_{116}N_{18}O_{20}$ by HR ESIMS. The NMR data for 5 were consistent with those of callipeltin C, except for the presence of a signal relative at a methyl group downfield shifted to $\delta = 1.79$ and of an olefinic signal at δ =6.70. One 2-amino-2-butenoic acid (dAbu) unit was assigned from the TOCSY and HMBC data, which provided correlations from the methyl signal at $\delta_{\rm H}$ = 1.79 to the olefinic carbon signals at δ 134.4 and 131.0 and from the olefinic proton to carbonyl signal a δ 166.6. ROESY correlations (in $CD₃OH$) between the –NH signal at $\delta = 9.74$ and the methyl signal at 1.79 indicated the (Z)-geometry for dAbu unit. The 2-amino-2-butenoic unit was found as common amino acid component of cyanobacteria hepatotoxins mycrocystins, $21-23$ where is present in both geometrical isomers, whereas in the sponge peptides aciculitins it was found as (E) isomer.²

Comparison of the reported NMR data of the two geometrical isomers of the dAbu unit allowed a

Table 1. ¹H and ¹³C NMR data (500 MHz, CD₃OD) for compounds **3**, **4** and **6**

aa	$\mathbf{3}$			4		6	
	$\delta_H{}^a$	$\delta_{\rm C}$	$\delta_{\rm H}$	$\delta_{\rm C}$		$\delta_{\rm H}$	$\delta_{\rm C}$
MeGlu					MeGlu		
α			5.14	57.6	α		
β			2.36, 2.02m	25.1	β		
Υ			2.20 ovl	32.9	Υ		
NMe			3.08	32.1	NMe		
$\rm CO$				173.8	$_{\rm CO}$		
COMH ₂				178.2	COMH ₂		
Leu					Leu		
α			4.84dd (9.7, 3.1)	49.7	α		
β			1.72, 1.52m	40.8	β		
Υ			1.70m	25.7	γ		
Me- γ			0.98d	23.4	$Me-$		
Me- ψ			0.96d	21.6	Me- ψ		
CONH				172.7	CONH		
Arg			4.38 ovl		Arg	4.40m	
α	4.48t(7.8)	54.2		53.9	α		58.0
β	1.97, 1.68m	29.2	1.93, 1.68m	29.6	β	1.96, 1.80m	29.6
$_{\delta}^{\gamma}$	1.60m	25.2	1.67m	26.4	γ	1.68m	26.4
	3.24m	41.9	3.19m	41.9	δ	3.22m	41.9
CONH		173.0		173.2	CONH		173.0
Guan		157.8		157.4	Guan		157.8
AlloThr-1					AlloThr		
α	4.33 ovl	59.4	4.36 ovl	60.5	α	4.31 ovl d	60.1
$\boldsymbol{\beta}$	4.13m	67.4	4.12m	68.6	β	4.17m	68.0
Υ	1.27d(7.0)	18.7	1.27d(7.0)	20.4	γ	1.29d(7.0)	20.1
CONH		171.7		172.7	CONH		172.1
AlloThr-2					dAbu		
α	4.33 ovl	59.7	4.22d(6.1)	61.2	α		130.0
β	4.09 ovl	67.4	4.05m	68.4	β	6.71q(7.9)	133.1
Υ	1.26d(6.0)	20.1	1.31d(6.0)	20.4	Υ	1.78d(7.9)	13.2
CONH		171.5		171.5	CONH		171.5
DiMeGln					DiMeGln		
α	4.44 ovl	57.6	4.45d(9.7)	57.8	α	4.32 ovl	58.0
β	2.23m	39.1	2.21 ovl	39.6	β	2.30 _m	37.1
β Me	1.02d(6.8)	14.0	1.01d(6.8)	13.8	β Me	1.13d(7.4)	14.0
	2.63 _m	41.8	2.63m	42.0	Υ	2.67m	42.6
Υ γ Me	1.20d(7.0)	19.9	1.18d(7.0)	15.8	γ Me	1.25d(7.4)	15.0
CONH		174.1		175.4	CONH		171.9
CONH ₂		179.3		179.2	CONH ₂		180.0
AGDHE					AGDHE		
α	3.98d(7.1)	73.4	3.98d(7.1)	72.9	α	4.02d(7.1)	73.4
β	3.70dd(7.1, 2.7)	75.0	3.76dd(7.1, 2.7)	75.3	β	3.80dd (7.1, 2.1)	74.0
$_{\delta}^{\gamma}$	4.15 ovl	50.2	4.15d(2.7)	50.7	γ	4.08m	51.0
	1.67, 1.30m	26.4	1.68, 1.30m	29.8	δ	1.67, 1.30m	29.6
$\boldsymbol{\varepsilon}$	1.67m	25.7	1.65m	25.4	$\boldsymbol{\varepsilon}$	1.67m	26.4
٣	3.25m	42.0	3.19m	41.9	Y	3.19m	42.0
$_{\rm CO}$		175.0		175.8	CO		176.1
Guan		157.8		157.8	Guan		157.8
Ala					Ala		
α	4.32 ovl	51.3	4.31q(7.3)	51.9	α	4.32 ovl	50.1
β	1.40d(7.3)	17.6	1.42d(7.3)	17.6	β	1.42d(7.6)	17.6
CONH		175.0		176.0	CONH		174.8
TMHEA					TMHEA		
1		177.9		178.8	$\mathbf{1}$		177.9
$\overline{\mathbf{c}}$	2.63m	44.9	2.62m	44.9	$\sqrt{2}$	2.63m	44.5
3	3.50d(d(8.7, 2.7))	78.9	3.51dd(8.7, 2.7)	76.5	3	3.50dd (8.7, 2.9)	78.9
4	1.77m	34.0	1.76m	33.7	4	1.75m	32.6
5	1.26m	40.4	1.23m	39.8	5	1.20m	38.4
6	1.69m	24.8	1.69m	26.5	6	1.69m	25.7
7	0.97d(6.2)	23.8	0.95d(6.2)	24.5	7	0.97d(6.7)	23.7
8	1.08d(6.0)	14.4	1.08d(6.0)	14.6	8	1.08d(6.9)	14.4
9	1.01d(6.5)	17.5	0.99d(6.5)	17.4	9	0.99d(6.9)	17.3
10	0.88d(6.5)	21.8	0.88d(6.5)	21.6	10	0.88d(6.7)	20.9

 $^{\rm a}$ Coupling constants are in parentheses and given in Hz. $^{\rm l}$ H and $^{\rm l}$ 3C assignments aided by COSY, TOCSY, HMQC and HMBC experiments. ovl: signal overlapped.

Figure 1. Callipeltin F (3) with ES MS/MS fragmentations.

confirmation of the proposed stereochemistry. The vinyl proton was observed at δ_H 6.5 ppm in the (Z)-isomer and at $\delta_{\rm H}$ 5.7 ppm in the (E)-isomer. Similarly the C-3 carbon is downfield shifted in the (Z)-isomer (δ _C 129.3 vs 123.8 ppm). NMR chemical shifts of the dAbu unit in callipeltin H [\(Table 2\)](#page-66-0) well mach those reported for the (Z)-isomer. It has been suggested that the dAbu residue could biogenetically arise by anti-dehydradation of the threonine^{[23](#page-69-0)} affording the E isomer from alloThr end the Z isomer from Thr. The co-occurrence in the same sponge of callipeltins containing D-*allo*Thr residue and of the corresponding derivatives containing the (Z)-dAbu residue rules out the hypothesis of a direct origin of callipeltin H and I from callipeltin C and F, respectively.

The amino acid sequence of 4 and placement of the acyl moiety were assigned from a combination of ESI MS/MS data, interesidue NOE interactions and HMBC correlations ([Table 2](#page-66-0) and [Figure 3\)](#page-67-0). The alignment of amino acid residues from the C-terminus N-MeAla to alloThr was the same as that of callipeltin C as secured by diagnostic b type fragmentations depicted in [Figure 3.](#page-67-0)

ROESY cross-peak between the *allo*Thr-NH (δ 7.86) and the dAbu-NH (δ 9.74) established an amide linkage between these two residues. The sequence of remaining units in callipeltin H was deduced from inter-residue NOE interaction NH/CH α as shown in [Table 2](#page-66-0).

The absolute configuration of the amino acid residues was

determined by Marfey's method to be D-alloThr, D-Arg, L-Leu, L-MeGlu, D-Ala.

To define the chirality of β -OMeTyr residue, a sample of 4 was ozonised and hydrolysed and subjected to Marfey's analysis. Ion selective monitoring for L-FDAA-OMeAsp $(m/z 416)$ showed a peak at $t_R 17.30$ corresponding to $2R,3S$ β -OMeAsp, indicating a 2R,3R stereochemistry of β -OMeTyr residue.^{[19](#page-69-0)}

The molecular formula $C_{42}H_{77}N_{13}O_{13}$ of callipeltin I (6), deduced by HR ESIMS, indicated the loss of one water molecule from callipeltin F. The presence of a 2-amino-2 butenoic acid (dAbu) unit was clearly inferred by the inspection of the ${}^{1}H$ NMR spectrum. The sequencing of the amino acid units was determined by the analysis of ESI MS/ MS fragmentation peaks [\(Fig. 4](#page-67-0)). The stereochemistry of the amino acid residues was determined by Marfey's method to be D-Ala, D-alloThr, D-Arg.

The finding of D -alloThr in all new callipeltins is in contrast with our previous studies on callipeltin $\overline{A}-E^3$ $\overline{A}-E^3$ and this results prompted us to reinvestigate the stereochemistry of the threonine residues in callipeltin A. Thus, callipeltin A was subjected to acid hydrolysis, Marfey's derivatisation and LC–MS analysis. A single peak corresponding to DalloThr was observed by ion-selective monitoring for FDAA-Thr (m/z 372). Our original misassignment (L-Thr vs D-alloThr) was due likely to the presence of an unassigned peak, 3 probably arising from a side reaction of a residue in callipeltin A during the hydrolysis. This fortuitously had the same retention time of FDAA derivative of L-Thr (12.5 min), but different molecular weight (m/z) 292).

The new callipeltin derivatives were tested for the anti HIV1 activity measured on infected human T-lymphoblasoid cells. No inhibition was observed even at high tested concentration of $2 \mu g/mL$, indicating the importance of the conformationally constrained cyclic structure for the antiviral activity observed for callipeltin A and related cyclodepsipetides.

Callipeltins F–I inhibit the growth of Candida albicans (ATCC24433) in the standard disk assay with a MIC of 10^{-4} M.

Figure 2. Callipeltin G (4) with ES MS/MS fragmentations.

(continued on next page)

Residue	$\delta_{\rm H}^{\rm \,a}$	$\delta_{\rm C}$	HMBC (1 H to 13 C)	NOE ^b
C ₅	7.21d(6.8)	129.7	C7	
C ₆	6.78d(6.8)	115.9		
C7		159.4		
OH	na			
OMe	3.14s	56.7	C ₂	
CONH	7.67 ovl	175.8		
MeAla				
α	5.12q(6.7)	57.7		
β	1.41d(6.7)	14.4	COOH	
NMe	2.84s	31.4	Cα, CO-βOMeTyr	

Table 2 (continued)

^a Coupling constants are in parentheses and given in Hz.

COOH 170.7

^b Correlations were obtained by NOESY with a 400 ms mixing time. na: not assigned. ovl: signal overlapped.

Figure 3. Callipeltin H (5) with ES MS/MS fragmentations.

Figure 4. Callipeltin I (6) with ES MS/MS fragmentations.

3. Experimental

3.1. General experimental procedures

Specific rotations were measured on a Perkin-Elmer 243 B polarimeter. High-resolution ESI-MS spectra were performed with a Micromass QTOF Micro mass spectrometer. ESI MS experiments were performed on a Applied Biosystem API 2000 triple-quadrupole mass spectrometer. NMR spectra were obtained on a Varian Mercury-400 and Inova- 500 NMR spectrometers (1 H at 400 and 500 MHz, 13 C at 100 and 125 MHz, respectively) equipped with a Bruker X-32 hardware, δ (ppm), J in Hz, spectra referred to CD_2HOD as internal standards (δ_{H} =3.30). HPLC was performed using a Waters Model 6000 A pump equipped with U6K injector and a differential refractometer, model 401.

3.2. Sponge material and separation of individual peptides

Latrunculia sp. (Family Latrunculidae, Demospongiae: Poecilosclerida) was collected at a depth of 15–20 m at Emae, Vanuatu South Pacific, in June 1996. The samples were frozen immediately after collection and lyophilised to yield 800 g of dry mass. Taxonomic identification was performed by Prof. John Hooper of Queensland Museum, Brisbane, Australia and reference specimens are on file (R1642) at the ORSTOM Centre of Noumea. Preliminary tests of bioactivity on polar extracts showed antifungal activity against Candida albicans and cytotoxic activity against L16 cells (10 µg/mL, 100% inhibition).

The lyophilised material (800 g) was extracted with methanol $(4 \times 2.5 \text{ L})$ at room temperature and the crude methanolic extract (80 g) was subjected to a modified Kupchan's partitioning procedure as follows. The methanol extract was dissolved in a mixture of MeOH/H₂O containing 10% H₂O and partitioned against *n*-hexane. The water content ($\%$ v/v) of the MeOH extract was adjusted to 20 and 40%, and partitioned against CCl_4 and CHCl_3 , respectively. The aqueous phase was concentrated to remove MeOH and then extracted with n-BuOH. The buthanol-soluble material (ca. 4 g) was chromatographated by DCCC in five runs $(CHCl₃/MeOH/H₂O, 7:13:8, ascending mode)$ and fractions of 4 mL were collected.

Fractions 10–11 were purified by HPLC on a Vydac C18 column (10 μ , 250 \times 10 mm, 4 mL/min) with CH₃CN/H₂O 28% (0.1% TFA) as eluent to give 4.8 mg of pure callipeltin F 3 (t_R =5.0 min). The additional peak at t_R =5.2 min was further purified by HPLC on a Thermo-Hypurity column (5 μ , 250 × 4.6 mm) eluting with CH₃CN/H₂O 24% containing 0.01% TFA (flow rate 1.2 mL/min) to give 2.4 mg of pure callipeltin I 6 (t_R =3.0 min).

Fractions 12–13 were purified by HPLC on a Vydac C18 column (10 μ , 250 \times 10 mm, 4 mL/min) with CH₃CN/H₂O 28% (0.1% TFA) as eluent to give a peak at t_R =5.0 min containing mainly callipeltin G (4) that was further purified by HPLC on a Thermo-Hypurity column $(5 \mu, 250 \times$ 4.6 mm) eluting with CH_3CN/H_2O 27% containing 0.05% TFA (flow rate 1.2 mL/min) to give 4.9 mg of pure 4 (t_R = 4.0 min).

Fractions 14–15 were purified by HPLC on a Vydac C18 column (10 μ , 250 × 10 mm, 4 mL/min) eluting with CH₃CN/H₂O 30% (0.1% TFA) to give a peak at t_R = 10.8 min containing a mixture of callipeltin C (2) and callipeltin H. Further purification by HPLC on a Thermo-Hypurity column (5 μ , 250 \times 4.6 mm) eluting with CH₃CN/ H2O 27% containing 0.05% TFA (flow rate 1.2 mL/min) gave 9.3 mg of pure callipeltin H 5 (t_R =6.2 min).

3.3. Characteristic of each compounds

3.3.1. Callipeltin F (3). 4.8 mg, white amorphous solid; $[\alpha]_D^{25}$ –4.3 (c 0.35, methanol); ¹H and ¹³C NMR data in CD3OD given in [Table 1;](#page-64-0) ESI-MS: m/z (%) 990.6 (35) $[M+H]^{+}$, 496.0 (100) $[M+2H]^{++}$. HRMS (ESI): calcd for C₄₂H₈₀N₁₃O₁₄: 990.5942; found 990.5938 [M+H]⁺.

3.3.2. Callipeltin G (4). 4.9 mg, white amorphous solid; $[\alpha]_D^{25}$ – 5.3 (c 0.26, methanol); ¹H and ¹³C NMR data in CD3OD given in [Table 1.](#page-64-0) ESI-MS: m/z (%) 1245.7 (100) $[M+H]$ ⁺. HRMS (ESI): calcd for $C_{54}H_{101}N_{16}O_{17}$: 1245.7525; found 1245.7532 $[M+H]$ ⁺.

3.3.3. Callipeltin H (5). 9.3 mg, white amorphous solid; $[\alpha]_D^{25}$ -4.5 (c 0.71, methanol); ¹H and ¹³C NMR data in CD3OH given in [Table 2.](#page-66-0) ESI-MS: m/z (%) 1505.7 (25) $[M+H]$ ⁺, 753.7 (100) $[M+2H]$ ⁺⁺. HRMS (ESI): calcd for $C_{68}H_{117}N_{18}O_{20}$: 1505.8686; found 1505.8677 [M+ $H]$ ⁺.

3.3.4. Callipeltin I (6). 2.4 mg, white amorphous solid; $[\alpha]_D^{25}$ 1.3 (c 0.37, methanol); ¹H and ¹³C NMR data in CD3OD given in [Table 1.](#page-64-0) ESI-MS: m/z (%) 972.5 (100) $[M+H]^+$. HRMS (ESI): calcd for $C_{42}H_{78}N_{13}O_{13}$: 972.5837; found 972.5848 $[M+H]$ ⁺.

3.4. Determination of absolute stereochemistry

3.4.1. General procedure for peptide hydrolysis. Peptide samples $(200 \mu g)$ were dissolved in degassed 6 N HCl $(0.5$ mL) in an evacuated glass tube and heated at 160 \degree C for 16 h. The solvent was removed in vacuo and the resulting material was subjected to further derivatisation.

3.4.2. General procedures for LC–MS analysis of Marfey's (FDAA) derivatives. A portion of the hydrolysate mixture (800 μ g) or the aminoacid standard (500 μ g) was dissolved in 80 μ L of a 2:3 solution of TEA/MeCN and treated with $75 \mu L$ of 1% 1-fluoro-2,4-dinitrophenyl-5-Lalaninamide (FDAA) in 1:2 MeCN/acetone. The vials were heated at 70° C for 1 h, and the contents were neutralised with 0.2 N HCl (50 μ L) after cooling to room temperature. An aliquot of the L-FDAA derivative was dried under vacuum, diluted with MeCN/5% HCOOH in H2O (1:1), and separated on a Vydac C18 $(25 \times 1.8 \text{ mm } \text{i.d.})$ column by means a linear gradient from 10 to 50% aqueous acetonitrile containing 5% formic acid and 0.05% trifluoracetic acid, over 45 min at 1 mL/min. The RP-HPLC system was connected to the electrospray ion source by inserting a splitter valve and the flow going into the mass spectrometer source was set at a value of $100 \mu L/min$. Mass spectra were acquired in positive ion detection mode (m/z interval of 320–900) and the data were analyzed using the suite of programs Xcalibur (ThermoQuest, San Jose´, California); all masses were reported as average values. Capillary temperature was set at 280° C, capillary voltage at 37 V, tube lens offset at 50 V and ion spray voltage at 5 V.

Retention times of authentic FDAA-amino acids (min): L-Thr (12.5) , D-Thr (17.6) , L-aThr (13.1) , D-aThr (14.1) , L-Ala (16.6 min), D-Ala (20.0 min), L-NMeAla (18.7 min), D-NMeAla (19.4 min), L-Arg (11.7 min), D-Arg (13.08 min), L-Leu (28.9 min), D-Leu (34.8 min).

The hydrolysate of callipeltin A contained: D-Arg (13.1), D-aThr (14.0), (3S,4R)-3,4-diMe-L-Glu (17.7), L-NMeAla (18.5), D-Ala (20.1), L-Leu (28.7).

The hydrolysate of callipeltin F contained: D-Arg (13.2), D-aThr (14.5), (3S,4R)-3,4-diMe-L-Glu (17.8).

The hydrolysate of callipeltin G contained: D-Arg (13.0), D-aThr (14.5), (3S,4R)-3,4-diMe-L-Glu (18.0), D-Ala (20.5) L-Leu (29.3).

The hydrolysate of callipeltin H contained: D-Arg (13.1), D-aThr (14.0), (3S,4R)-3,4-diMe-L-Glu (18.0), D-Ala (20.2), L-NMeAla (18.5 min), L-Leu (29.7).

The hydrolysate of callipeltin I contained: D-Arg (13.4), D-aThr (14.2), (3S,4R)-3,4-diMe-L-Glu (18.5).

3.5. Determination of the absolute stereochemistry of β -OMeTyr residue in callipeltin H (5)

A stream of ozone in O_2 was bubbled through a cooled solution of callipeltin H (0.5 mg) or of all four diastereomers of β -OMeTyr (1 mg)^{[19](#page-69-0)} in MeOH (0.5 mL) at -78 °C for 1 h. Hydrogen peroxide (35%, 10 drops) was added to the reaction mixture with then allowed to stand at room temperature overnight. The solvent was removed under a stream of N_2 and the resulting β -methoxyaspartates were immediately subjected to Marfey derivatization. The ozonolysis product of callipeltin H was then dissolved in

degassed 6 N HCl (0.5 mL) in an evacuated glass tube and heated at 160° C for 16 h. The solvent was removed in vacuo and the resulting material was subjected to further derivatisation.

A portion of callipeltin H hydrolizate mixture or the β -methoxyaspartates (500 µg) was subjected to Marfey's derivatization and LC–MS analysis.

Retention times of authentic L-FDAA-β-OMeAsps (min): $(2S,3S)$ - β -OMeAsp (10.23 min) , $(2S,3R)$ - β -OMeAsp (16.06 min), $(2R,3R)$ - β -OMeAsp (9.86 min) , $(2R,3S)$ - β -OMeAsp (17.30 min). The hydrolysate of ozonolysis product of callipeltin H contained: $(2R,3S)$ - β -OMeAsp (17.30 min).

3.6. Antifungal tests

The broth macrodilution test was performed by using the NCCLS standard reference method for broth dilution antifungal susceptibility testing of yeasts. 25 Stock solutions of callipeltins $\overline{F}-I$ (10⁻³ M) were prepared, divided into aliquots, and stored at -80 °C. A new aliquot was thawed on each day of use. Before testing, Candida albicans (ATCC 24433) were maintained on Sabouraud's agar slants and periodically transferred to Sabouraud's agar plates and incubated for 48 h at 28° C. To prepare stationary growth phase yeast, a colony was taken from the agar plate and transferred into 30 mL Sabouraud-dextrose broth (DIFCO laboratories, Detroit, MI) and incubated for 72 h at 35 $^{\circ}$ C. Cells were centrifuged at $1000 \times g$ for 10 min and the pellet was washed twice with distilled water. Cells were counted and suspended in RPMI 1640 plus 0.165 M MOPS buffer at a density of 5000 CFU/mL. One hundred microlitres of the yeast suspension was transferred into control wells or wells containing the callipeltins 10^{-4} – 10^{-8} M final concentrations. The plates were incubated in air at 35° C without agitation for 48 h. The experiments were run in triplicate. Negative growth corresponded to no visible growth in the well.

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A concise synthesis of polyhydroxydihydrochalcones and homoisoflavonoids

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Abstract—A general and single step synthesis of polyhydroxydihydrochalcones from the readily available phenols and dihydrocinnamic acids using BF_3 Et₂O is described. The method allows the synthesis of a wide range of compounds with multiple phenolic hydroxyls and other substituents. These dihydrochalcones are converted into homoisoflavonoids by DMF/PCl5 and the methodology has been applied to the synthesis of naturally occurring phloretin and 5,7-dihydroxy-3-[(4-hydroxyphenyl)methyl]-4H-chromen-4-one. The antioxidant activity of dihydrochalcones and homoisoflavonoids was determined by superoxide free radical (NBT) and DPPH free radical scavenging methods. Polyhydroxydihydrochalcones 3c, 3f, 3g and homoisoflavonoids 4c, 4f, 4g displayed excellent antioxidant activity. $© 2005 Elsevier Ltd. All rights reserved.$

1. Introduction

Naturally occurring polyphenolic compounds display wide spectrum of biological activities. Polyphenolic flavonoids have gained increasing importance in view of their strong antioxidant activity and preventing role in free radical mediated disorders such as cancer, Alzheimer's, Parkinson's, and cardiovascular diseases.^{[1](#page-74-0)} Dihydrochalcones (DHCs), the reduced form of chalcones, have been known to occur widely in nature and are important intermediates for many natural products and pharmaceutical drugs. $¹$ $¹$ $¹$ DHCs have been reported to have various</sup> biological activities $2,3$ and have received considerable attention as food sweeteners (neohesperidin).^{[4](#page-74-0)} Up to now, DHCs were mostly synthesized through a Claisen–Schmidt condensation^{[5,6](#page-74-0)} to obtain chalcone, followed by reduction to DHC or through the alkaline reduction of a correspon-ding flavanone.^{[7,8](#page-74-0)} Although widely used, the procedures are not suitable for polyhydroxyDHCs, as these methods require protection of all phenolic hydroxyls. Recently, palladium mediated coupling of iodobenzenes and the enol of acetophenone was reported σ as an alternative route to DHCs. The homoisoflavonoids (3-benzylchromen-4-ones), are naturally occurring compounds and are structurally related to flavonoids,^{[10](#page-74-0)} and display a wide spectrum of biological activities.^{[11–15](#page-74-0)} A few methods of synthesis have been reported in the literature for homoisoflavonoids and these were based on (i) the condensation of 4-chromanones with arylaldehydes in methanol by passing HCl gas or by using piperidine as a base^{[16,17](#page-75-0)} followed by isomerisation of the double bond using Pd/C at 250° C, (ii) hydrogenation of chalcones followed by one carbon extension using ethyl formate/sodium^{[18](#page-75-0)} or methanesulfonyl chloride/ \overline{DMF} . Both the methods have disadvantages; while the first method has multiple steps, in the second method, the phenolic hydroxyls have to be protected to get chalcones in good yield. Our interest in the chemistry of the flavonoids^{[20](#page-75-0)} and an increasing demand for a short, and efficient method prompted us to develop a simple and general method for the synthesis of DHCs and homoisoflavonoids. The methodology has been applied to the synthesis of phloretin, $2¹$ a naturally occurring DHC and $5,7$ -dihydroxy-3-[(4-hydroxyphenyl)-methyl]- $4H$ -chromen-4-one, a homoisoflavonoid from Ophiopogon jaburan.^{[22](#page-75-0)} Moreover, to the best of our knowledge, there is no report in the literature on the antioxidant activity of homoisoflavonoids. So we report in this paper, the details of synthesis of DHCs, homoisoflavonoids and their antioxidative activity results.

Keywords: Dihydrochalcones; Homoisoflavonoids; Boron trifluoride etherate; Antioxidant activity.

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2. Results and discussion

2.1. Synthesis

During the course of our investigations of new synthetic routes to flavonoids, we have found that a Friedel-Crafts reaction constitutes a novel and efficient approach to DHCs. The method involves the preparation of DHCs in a single step from phenols and dihydrocinnamic acids by Friedel-Crafts acylation using boron trifluoride etherate, which serves as the Lewis acid for the acylation and as solvent for the reaction (Scheme 1). In this method, protection of the phenolic hydroxyls is not necessary. In a typical experiment, substituted phenol (1, 3 mmol) and dihydrocinnamic acid (2, 3 mmol) was treated with boron trifluoride etherate at 80–90 \degree C for 90 min. After completion of the reaction (monitored by TLC), it was poured into aqueous sodium acetate and extracted with ethyl acetate to give DHCs (3). The generality of the reaction was established with various phenols and substituted dihydrocinnamic acids and in all cases (Table 1, 7 examples) the reaction was completed within 90 min. In the second step, the DHCs $(3a-3g)$ were treated with N, N' -dimethyl(chloromethylene)ammonium chloride^{[23](#page-75-0)} generated in situ from DMF and PCl₅ for one carbon extension to get homoisoflavonoids $(4a-4g)$ in 78–88% yield (Table 1). In all cases the reaction was complete in 2 h and the products were characterized by their spectral data (IR, NMR and Mass). The methodology has been applied to the synthesis of naturally occurring phloretin,[21](#page-75-0) a DHC and 5,7-dihydroxy-3-[(4-hydroxyphenyl)methyl]-4H-chromen-4-one (4h), isolated from Ω *phiopogon jaburan*.^{[22](#page-75-0)} Reaction of phloroglucinol and 4-hydroxydihydrocinnamic acid with $BF_3 \cdot Et_2O$ gave phloretin (3h), which was converted further into 4h (85% y ield) using N, N' -dimethyl(chloromethylene)ammonium

chloride. The spectral data of synthetic 3h and 4h were found to be identical with those of the corresponding natural products.^{[19](#page-75-0)}

2.2. Antioxidant activity

We have determined the antioxidative activity of DHCs (3a–3h) and homoisoflavonoids (4a–4h) by nitro blue tetrazolium $(NBT)^{24,25}$ $(NBT)^{24,25}$ $(NBT)^{24,25}$ and 1,1-diphenyl-2-picrylhydrazyl $(DPPH)²⁶$ $(DPPH)²⁶$ $(DPPH)²⁶$ free radical scavenging methods. The $IC₅₀$ values of these compounds are presented in [Table 2.](#page-72-0) DHCs 3f $(IC_{50}: 12.9 \mu M),$ 3g $(IC_{50}: 19.6 \mu M)$ and 3c $(IC_{50}: 30.2 \mu M)$ and homoisoflavonoids $4f$ (IC₅₀: 6.3 µM), $4g$ (IC₅₀: 8.2 µM) and $4c$ (IC₅₀: 24.8 μ M) having catechol moieties were the most active compounds. Interestingly 3f, 3g and 3c and 4f, 4g and 4c showed several-fold more potent activity than vitamin C (IC₅₀: 852 µM), vitamin E (IC₅₀: 726 µM), BHA $(IC_{50}: 966 \mu M)$ and BHT $(IC_{50}: 381 \mu M)$. The same order of activity was followed by DHCs 3a–3h and homoisoflavonoids 4a–4h with the DPPH method. Again 3c, 3f and 3g and 4c, 4f and 4g showed good DPPH free radical scavenging activity. The superior antioxidative activity of these compounds lends further support to the fact that the catechol system enhances the antioxidative activity.^{[27](#page-75-0)}

3. Conclusions

In conclusion, we have described a general, single step method for the synthesis of polyhydroxydihydrochalcones using phenols and dihydrocinamic acids with $BF_3 \cdot Et_2O$. One carbon extension of these dihydrochalcones into homoisoflavonoids by $DMF/PCl₅$ was achieved in good yields. The DHCs and homoisoflavonoids were evaluated for their antioxidative potential by two commonly used

Scheme 1. Reagents and conditions: (i) $BF_3 \cdot Et_2O$, 80–90 °C, 90 min, 30–71% (ii) $BF_3 \cdot Et_2O$, DMF/PCl₅, rt, 2 h, 78–88%.

Table 1. Dihydrochalcones 3 and homoisoflavonoids 4

S.no.	Entry	ĸ۱	R_{2}	R_3	R_4	R_5	K_6	3 Yield $(\%)^a$	4 Yield $(\%)^a$
			OH		Н		OН	62	85
			OН		Н	Н	OCH ₃		88
			OH			ΟH	OН	55	82
			OН		Н	OCH ₃	OCH ₃	68	87
	Δ		OΗ		OCH ₃	Н	OCH ₃	61	78
6		OН	OH		Н	H	OН	58	80
		OН	OΗ			Н	OCH ₃	65	81
			OΗ	OН	Н		OН	30	85

^a Unoptimized isolated yields.
Table 2. Antioxidant activity of homoisoflavonoids

Compound no.	NBT superoxide scavenging activity (IC ₅₀ in μ M)	DPPH free radical scaven- ging activity (IC_{50} in μ M)
3a	>100	>100
3 _b	>100	>100
3c	30.2	7.8
3d	>100	>100
3e	>100	>100
3f	12.9	11.3
3 _g	19.6	14.9
3h	16.6	95
4a	>100	>100
4b	>100	>100
4c	24.8	13.4
4d	>100	>100
4e	>100	>100
4f	6.3	19.0
4g	8.2	19.6
4h	17.6	>100
Vitamin C	852	25.1
Vitamin E	726	>100
BHA	966	34.0
BHT	381	22.5

BHA, butylated hydroxyanisole; BHT, butylated hydroxytoluene; NBT, nitro blue tetrazolium; DPPH, 1,1-diphenyl-2-picrylhydrazyl. The lower the IC ϵ_0 values, the higher is the antioxidant activity.

methods, the superoxide and DPPH free radical scavenging methods. DHCs 3c, 3f and 3g and homoisoflavonoids 4c, 4f and 4g were potent antioxidants.

4. Experimental

4.1. General

Melting points were recorded on a Mel-Temp melting point apparatus, in open capillaries and are uncorrected. IR spectra were recorded on a Perkin-Elmer BX1 FTIR Spectrophotometer and ${}^{1}H$ NMR (400 MHz), ${}^{13}C$ NMR-DEPT (100 MHz) spectra were recorded on a Bruker AMX 400 MHz NMR spectrometer using TMS as internal standard and the values for chemical shifts (δ) being given in ppm and coupling constants (J) in Hertz (Hz) . Mass spectra were recorded on an Agilent 1100 LC/MSD. Acme silica gel G and silica gel (100–200 mesh) were used for analytical TLC and column chromatography, respectively.

4.2. General procedure for dihydrochalcones (3)

A mixture of phenol (1, 3 mmol), 3-phenylpropanoic acid $(2, 3 \text{ mmol})$ and $BF_3 \text{·} Et_2O$ $(1.94 \text{ mL}, 15.3 \text{ mmol})$ was stirred at 80–90 °C for 90 min under N_2 . The reaction mixture was poured into 10% aqueous NaOAc solution (100 mL) and allowed to stand for 4 h and the solution was extracted with EtOAc $(3 \times 100 \text{ mL})$. The combined EtOAc layer was washed with water (20 mL), brine (20 mL) and dried over $Na₂SO₄$. The residue obtained after evaporation of the solvent was chromatographed over silica gel column using hexane–EtOAc mixtures as eluent to give 3a–h.

4.2.1. 1-(2,4-Dihydroxyphenyl)-3-(4-hydroxyphenyl) propan-1-one (3a). Light brown powder (480 mg, 62%), mp 140–142 °C; IR (KBr): 3456, 3270, 1626, 1214, 1165, 1134, 987 cm⁻¹; ¹H NMR (DMSO-d₆): δ 12.63 (1H, s, Ar-OH), 11.69 (1H, s, Ar-OH), 10.60 (1H, s, Ar-OH), 7.79

 $(1H, d, J=8.8 \text{ Hz}, H-6^{\prime\prime}), 7.04 (2H, d, J=8.3 \text{ Hz}, H-2^{\prime}, 6^{\prime}),$ 6.65 (2H, d, $J=8.3$ Hz, $H=3', 5'$), 6.35 (1H, dd, $J=8.8$) 2.4 Hz, H-5"), 6.24 (1H, d, $J=2.4$ Hz, H-3"), 3.20 (2H, t, $J=7.6$ Hz, H-3), 2.81 (2H, t, $J=7.6$ Hz, H-2); ¹³C NMR $(DMSO-d₆)$: δ 203.9, 164.7, 164.3, 155.5, 133.0, 131.0, 129.3, 115.1, 112.6, 108.2, 102.4, 39.1, 29.1; MS (ESI, negative ion mode): m/z 257 (M-H)⁻. Analysis found: C, 69.68; H, 5.52%. Calcd for $C_{15}H_{14}O_4$: C, 69.76; H, 5.46%.

4.2.2. 1-(2,4-Dihydroxyphenyl)-3-(4-methoxyphenyl) propan-1-one (3b). Light brown powder (580 mg, 71%), mp 58–60 °C; IR (KBr): 3456, 3105, 1629, 1225, 1131, $1029, 990 \text{ cm}^{-1}$; ¹H NMR (DMSO-d₆): δ 12.60 (1H, s, Ar-OH), 10.61 (1H, s, Ar-OH), 7.79 (1H, d, $J=8.8$ Hz, H-6ⁿ), 7.16 (2H, d, $J=8.3$ Hz, $H=2^{\prime},6^{\prime}$), 6.82 (2H, d, $J=$ 8.3 Hz, H-3',5'), 6.34 (1H, dd, $J=8.8$, 2.0 Hz, H-5"), 6.23 $(H, d, J=2.0 \text{ Hz}, H-3^{\prime\prime}), 3.69 \text{ (3H, s, Ar-OCH₃), 3.22 \text{ (2H,$ t, $J=7.5$ Hz, H-3), 2.84 (2H, t, $J=7.5$ Hz, H-2); ¹³C NMR (DMSO-d6): d 203.7, 164.8, 164.3, 157.6, 132.9, 129.4, 114.2, 113.7, 112.6, 108.2, 102.5, 55.0, 40.1, 29.0; MS (ESI, negative ion mode): m/z 271 (M-H)⁻. Analysis found: C, 70.52; H, 5.97%. Calcd for C₁₆H₁₆O₄: C, 70.58; H, 5.92%.

4.2.3. 1-(2,4-Dihydroxyphenyl)-3-(3,4-dihydroxyphenyl) propan-1-one (3c). Colorless powder (450 mg, 55%), mp 108–110 °C; IR (KBr): 3339, 1638, 1605, 1286, 1222, 1200, 1174, 1141, 968 cm⁻¹; ¹H NMR (DMSO- d_6): δ 12.63 (1H, s, Ar-OH), 8.72 (1H, s, Ar-OH), 8.56 (1H, s, Ar-OH), 8.29 (1H, s, Ar-OH), 7.77 (1H, d, $J=8.8$ Hz, H-6ⁿ), 7.10–7.20 (2H, m, $H-2^{\prime},5^{\prime}$), 6.47 (1H, dd, J=7.8, 1.5 Hz, H-6^{\prime}), 6.34 (1H, dd, $J=8.8$, 2.0 Hz, H-5^{$\prime\prime$}), 6.23 (1H, d, $J=$ 2.0 Hz, H-3"), 3.16 (2H, t, $J=7.6$ Hz, H-3), 2.74 (2H, t, $J=$ 7.6 Hz, H-2); ¹³C NMR (DMSO- d_6): δ 205.0, 165.9, 165.4, 146.1, 144.5, 134.1, 133.0, 120.1, 117.0, 116.6, 113.7, 109.3, 103.6, 40.4, 30.4; MS (ESI, negative ion mode): m/z 273 $(M-H)^{-}$. Analysis found: C, 65.64; H, 5.18%. Calcd for $C_{15}H_{14}O_5$: C, 65.69; H, 5.15%.

4.2.4. 1-(2,4-Dihydroxyphenyl)-3-(3,4-dimethoxyphenyl) propan-1-one (3d). Light brown powder (615 mg, 68%), mp 128–130 °C; IR (KBr): 3371, 1631, 1257, 1235, 1209, 1137, 1026, 991 cm⁻¹; ¹H NMR (DMSO-d₆): δ 12.64 (1H, s, Ar-OH), 10.63 (1H, s, Ar-OH), 7.82 (1H, d, $J=8.8$ Hz, H-6^{*i*}), 6.88 (1H, d, $J=1.8$ Hz, H-2^{*i*}), 6.83 $(1H, d, J=8.0 \text{ Hz}, H=5^{\prime}), 6.76 \text{ (1H, dd, } J=8.0, 1.8 \text{ Hz}, H=$ $6'$), 6.37 (1H, dd, $J=8.8$, 2.1 Hz, H-5ⁿ), 6.26 (1H, d, $J=$ 2.1 Hz, H-3"), 3.73 (3H, s, Ar-OCH₃), 3.70 (3H, s, Ar-OCH₃), 3.25 (2H, t, $J=7.6$ Hz, H-3), 2.86 (2H, t, $J=7.6$ Hz, H-2); ¹³C NMR (DMSO- d_6): δ 203.8, 164.8, 164.3, 148.7, 147.2, 133.5, 133.0, 120.2, 112.6, 112.4, 111.9, 108.2, 102.5, 55.5, 55.4, 39.2, 29.5; MS (ESI, negative ion mode): m/z 301 (M-H)⁻. Analysis found: C, 67.52; H, 6.04%. Calcd for $C_{17}H_{18}O_5$: C, 67.54; H, 6.00%.

4.2.5. 1-(2,4-Dihydroxyphenyl)-3-(2,4-dimethoxyphenyl) propan-1-one (3e). Colorless powder (555 mg, 61%), mp 110–112 °C; IR (Neat): 3347, 1626, 1290, 1208, 1153, 1037, 990 cm⁻¹; ¹H NMR (DMSO- d_6): δ 12.72 (1H, s, Ar-OH), 9.99 (1H, s, Ar-OH), 7.60 (1H, d, $J=8.5$ Hz, H-6^{$\prime\prime$}), 7.00 (1H, d, J=8.5 Hz, H-5^{$\prime\prime$}), 6.20–6.40 (4H, m, $H-3',5',6',3'')$, 3.81 (3H, s, Ar-OCH₃), 3.78 (3H, s, Ar-OCH₃), 3.10 (2H, t, $J=7.5$ Hz, H-3), 2.90 (2H, t, $J=7.5$ Hz, H-2); ¹³C NMR (DMSO-d₆): δ 204.2, 164.8,

164.4, 159.2, 158.0, 132.9, 130.0, 120.8, 112.5, 108.2, 104.3, 102.5, 98.3, 55.5, 55.3, 38.0, 24.8; MS (ESI, negative ion mode): m/z 301 (M – H)⁻. Analysis found: C, 67.51; H, 6.05%. Calcd for $C_{17}H_{18}O_5$: C, 67.54; H, 6.00%.

4.2.6. 1-(2,3,4-Trihydroxyphenyl)-3-(4-hydroxyphenyl) propan-1-one (3f). Pale green powder (475 mg, 58%), mp 158–160 8C; IR (KBr): 3440, 3246, 1633, 1243, 1213, 1118, 1044, 1004, 899 cm⁻¹; ¹H NMR (DMSO- d_6): δ 12.63 (1H, s, Ar-OH), 10.06 (1H, s, Ar-OH), 9.15 (1H, s, Ar-OH), 8.59 (1H, s, Ar-OH), 7.34 (1H, d, $J=8.8$ Hz, H-6ⁿ), 7.04 (2H, d, $J=8.3$ Hz, H-2',6'), 6.66 (2H, d, $J=8.3$ Hz, $H=3',5'$), 6.38 $(1H, d, J=8.8 \text{ Hz}, H=5^{\prime\prime}), 3.20 \ (2H, t, J=7.5 \text{ Hz}, H=3), 2.82$ (2H, t, J=7.5 Hz, H-2); ¹³C NMR (DMSO- d_6): δ 204.9, 155.7, 152.6, 152.4, 132.5, 131.2, 129.4, 122.6, 115.2, 113.0, 107.9, 39.0, 29.4; MS (ESI, negative ion mode): m/z 273 $(M-H)^{-}$. Analysis found: C, 65.66; H, 5.20%. Calcd for $C_{15}H_{14}O_5$: C, 65.69; H, 5.15%.

4.2.7. 1-(2,3,4-Trihydroxyphenyl)-3-(4-methoxyphenyl) **propan-1-one (3g).** Light brown oil $(560 \text{ mg}, 65\%)$, IR (Neat): 3396, 2928, 1632, 1244, 1178, 1114, 1033, 999 cm⁻¹;
¹H NMP (DMSO d): δ 12.60 (1H s, At OH), 10.08 (1H s, ¹H NMR (DMSO- d_6): δ 12.60 (1H, s, Ar-OH), 10.08 (1H, s, Ar-OH), 8.61 (1H, s, Ar-OH), 7.33 (1H, d, $J=8.8$ Hz, H-6ⁿ), 7.16 (2H, d, $J=7.8$ Hz, $H=2^{\prime},6^{\prime}$), 6.81 (2H, d, $J=7.8$ Hz, $H-3',5'$), 6.38 (1H, d, $J=8.8$ Hz, $H-5''$), 3.69 (3H, s, Ar-OCH₃), 3.20 (2H, t, $J=7.5$ Hz, H-3), 2.85 (2H, t, $J=$ 7.5 Hz, H-2); ¹³C NMR (DMSO- d_6): δ 204.8, 157.8, 152.7, 152.6, 133.1, 132.6, 129.6, 122.6, 114.0, 113.1, 108.0, 55.2, 39.4, 29.3; MS (ESI, negative ion mode): m/z 287 (M-H)⁻. Analysis found: C, 66.62; H, 5.62%. Calcd for $C_{16}H_{16}O_5$: C, 66.66; H, 5.59%.

4.2.8. 1-(2,4,6-Trihydroxyphenyl)-3-(4-hydroxyphenyl) propan-1-one (3h). Pale yellow solid (245 mg, 30%), mp 258–260 °C (lit.^{[19](#page-75-0)} mp 257–258 °C); IR (KBr): 3268, 1630, 1606, 1296, 1209, 1163, 1076 cm⁻¹; ¹H NMR (DMSO- d_6): d 12.23 (2H, s, Ar-OH), 10.36 (1H, s, Ar-OH), 9.13 (1H, s, Ar-OH), 7.00 (2H, d, $J=8.3$ Hz, H-2',6'), 6.65 (2H, d, $J=$ 8.3 Hz, H-3',5'), 5.79 (2H, s, H-3", 5"), 3.20 (2H, t, $J=$ 7.8 Hz, H-3), 2.74 (2H, t, $J=7.8$ Hz, H-2); ¹³C NMR (DMSO-d6): d 204.4, 164.8, 164.4, 155.6, 131.9, 129.4, 115.3, 103.9, 94.9, 45.7, 29.6; MS (ESI, negative ion mode): m/z 273 $(M-H)^{-}$.

4.3. General procedure for homoisoflavonoids (4)

A mixture of 3 (3 mmol) and $BF_3 \cdot Et_2O$ (1.2 mL, 9 mmol) was cooled to 10 \degree C and DMF (4.6 mL) was added drop wise for 5 min. In another flask, DMF (8 mL) was cooled to 10° C and PCl₅ (0.939 g, 4.5 mmol) was added in small portions. The mixture was then allowed to stand to 55 \degree C for 20 min. The light yellow colored solution containing N, N' dimethyl(chloromethylene)ammonium chloride was then added to the above reaction mixture slowly at $20-25$ °C. The mixture was stirred at rt for 2 h and poured into boiling dil HCl slowly and cooled. The solution was extracted with EtOAc $(3 \times 50 \text{ mL})$ and the combined EtOAc layer was washed with water (20 mL), brine (20 mL) and dried over sodium sulfate. The residue obtained after evaporation of the solvent was chromatographed over silica gel column using chloroform–methanol mixtures as eluent to give 4a–h.

4.3.1. 7-Hydroxy-3-[(4-hydroxyphenyl)methyl]-4Hchromen-4-one (4a). Colorless powder (680 mg, 85%), mp 212–214 °C; IR (KBr): 3430, 1628, 1600, 1267, 1244, 1176, 1140, 1099, 960 cm⁻¹; ¹H NMR (DMSO- d_6): δ 10.73 $(2H, br s, Ar-OH), 8.04 (1H, s, H-2), 7.86 (1H, d, J=8.8 Hz,$ $H=5$), 7.06 (2H, d, $J=8.3$ Hz, $H=2^{\prime},6^{\prime}$), 6.88 (1H, dd, $J=8.8$, 2.2 Hz, H-6), 6.79 (1H, d, $J=2.2$ Hz, H-8), 6.65 (1H, d, $J=$ 8.3 Hz, H-3['],5'), 3.53 (2H, s, H-9); ¹³C NMR (CDCl₃+ DMSO-d6): d 176.3, 162.2, 157.8, 155.4, 152.2, 129.4, 128.9, 126.6, 123.8, 116.3, 115.1, 114.6, 102.0, 30.2; MS (ESI, negative ion mode): m/z 267 $(M-H)^-$. Analysis found: C, 71.59; H, 4.55%. Calcd for C₁₆H₁₂O₄: C, 71.64; H, 4.51%.

4.3.2. 7-Hydroxy-3-[(4-methoxyphenyl)methyl]-4Hchromen-4-one (4b). Light brown powder (740 mg, 88%), mp 162–164 °C; IR (KBr): 3433, 1631, 1248, 1161, 1132, 1096, 1034 cm⁻¹; ¹H NMR (DMSO- d_6): δ 10.55 (1H, s, Ar-OH), 7.96 (1H, s, H-2), 7.87 (1H, d, $J=8.8$ Hz, H-5), 7.19 (2H, d, $J=8.3$ Hz, $H=2^{\prime},6^{\prime}$), 6.86 (1H, dd, $J=8.8$) 2.0 Hz, H-6), 6.80 (2H, d, $J=8.3$ Hz, H-3^{\prime},5^{\prime}), 6.71 (1H, d, $J=2.0$ Hz, H-8), 3.72 (3H, s, Ar-OCH₃), 3.61 (2H, s, H-9); ¹³C NMR (CDCl₃+DMSO- d_6): δ 176.2, 162.2, 157.8, 157.6, 152.1, 130.6, 129.4, 126.6, 123.5, 116.3, 114.6, 113.4, 102.0, 54.7, 30.2; MS (ESI, negative ion mode): m/z 281 $(M-H)^{-}$. Analysis found: C, 72.29; H, 5.04%. Calcd for C17H14O4: C, 72.33; H, 5.00%.

4.3.3. 7-Hydroxy-3-[(3,4-dihydroxyphenyl)methyl]-4Hchromen-4-one (4c). Colorless powder (695 mg, 82%), mp 192-194 °C; IR (KBr): 3393, 1627, 1239, 1180, 1113, 967 cm⁻¹; ¹H NMR (DMSO- d_6): δ 10.73 (1H, s, Ar-OH), 8.72 (1H, s, Ar-OH), 8.61 (1H, s, Ar-OH), 8.07 (1H, s, H-2), 7.86 (1H, d, $J=8.8$ Hz, H-5), 6.88 (1H, dd, $J=8.8$, 2.2 Hz, H-6), 6.79 (1H, d, $J=2.2$ Hz, H-8), 6.63 (1H, d, $J=2.1$ Hz, $H-2'$), 6.59 (1H, d, $J=8.3$ Hz, $H-5'$), 6.51 (1H, dd, $J=8.3$, 2.1 Hz, H-6'), 3.47 (2H, s, H-9); ¹³C NMR (CDCl₃+ DMSO-d6): d 176.4, 162.1, 157.8, 152.3, 144.4, 142.9, 129.9, 126.6, 123.7, 119.8, 116.3, 115.7, 115.1, 114.6, 102.0, 30.3; MS (ESI, negative ion mode): m/z 283 $(M-H)^-$. Analysis found: C, 67.58; H, 4.29%. Calcd for $C_{16}H_{12}O_5$: C, 67.60; H, 4.26%.

4.3.4. 7-Hydroxy-3-[(3,4-dimethoxyphenyl)methyl]-4Hchromen-4-one (4d). Colorless powder (815 mg, 87%), mp 180–182 °C; IR (KBr): 3245, 1632, 1262, 1244, 1175, 1140, 1094, 1026, 963 cm⁻¹; ¹H NMR (DMSO- d_6): δ 10.62 $(1H, s, Ar-OH), 8.01 (1H, s, H-2), 7.88 (1H, d, J=8.8 Hz,$ H-5), 6.78–6.90 (5H, m, Ar-H), 3.75 (3H, s, Ar-OCH3), 3.72 $(3H, s, Ar-OCH₃), 3.61 (2H, s, H-9);$ ¹³C NMR (CDCl₃+ DMSO-d6): d 176.0, 162.3, 157.8, 152.4, 148.4, 147.1, 131.4, 126.6, 123.3, 120.5, 116.3, 114.7, 112.2, 111.2, 102.0, 55.4, 55.3, 30.6; MS (ESI, negative ion mode): m/z 311 $(M-H)^{-}$. Analysis found: C, 69.19; H, 5.21%. Calcd for $C_{18}H_{16}O_5$: C, 69.22; H, 5.16%.

4.3.5. 7-Hydroxy-3-[(2,4-dimethoxyphenyl)methyl]-4Hchromen-4-one (4e). Colorless powder (730 mg, 78%), mp 182-184 °C; IR (KBr): 3224, 1631, 1241, 1207, 1158, 1124, 1040 cm⁻¹; ¹H NMR (DMSO-d₆): δ 10.72 (1H, s, Ar-OH), 7.86 (1H, d, $J=8.8$ Hz, H-5), 7.85 (1H, s, H-2), 7.02 (1H, d, $J=8.3$ Hz, H-6^{\prime}), 6.88 (1H, dd, $J=8.8$, 2.0 Hz, H-6), 6.79 (1H, d, $J=2.0$ Hz, H-8), 6.51 (1H, d, $J=2.1$ Hz,

H-3[']), 6.42 (1H, dd, $J=8.3$, 2.1 Hz, H-5[']), 3.77 (3H, s, Ar-OCH3), 3.70 (3H, s, Ar-OCH3), 3.51 (2H, s, H-9); MS (ESI, negative ion mode): m/z 311 (M-H)⁻. Analysis found: C, 69.18; H, 5.20%. Calcd for $C_{18}H_{16}O_5$: C, 69.22; H, 5.16%.

4.3.6. 7,8-Dihydroxy-3-[(4-hydroxyphenyl)methyl]-4Hchromen-4-one (4f). Colorless powder (680 mg, 80%), mp 251–253 °C; IR (KBr): 3458, 1628, 1201, 1175, 1156, $1047, 985 \text{ cm}^{-1}$; ¹H NMR (DMSO-d₆): δ 10.17 (1H, s, Ar-OH), 9.15 (2H, br s, $2 \times$ Ar-OH), 8.14 (1H, s, H-2), 7.37 $(1H, d, J=8.6 \text{ Hz}, H=5)$, 7.07 $(2H, d, J=8.3 \text{ Hz}, H=2', 6')$, 6.91 (1H, d, $J=8.6$ Hz, H-6), 6.65 (2H, d, $J=8.3$ Hz, H-3',5'), 3.56 (2H, s, H-9); ¹³C NMR (CDCl₃+DMSO-d₆): d 176.1, 155.6, 152.9, 149.9, 147.1, 132.9, 129.9, 129.5, 123.0, 117.1, 115.2, 115.0, 114.1, 30.0; MS (ESI, negative ion mode): m/z 283 (M – H)⁻. Analysis found: C, 67.57; H, 4.30%. Calcd for $C_{16}H_{12}O_5$: C, 67.60; H, 4.26%.

4.3.7. 7,8-Dihydroxy-3-[(4-methoxyphenyl)methyl]-4Hchromen-4-one (4g). Colorless powder (725 mg, 81%), mp 252–254 °C; IR (KBr): 3320, 3150, 1631, 1241, 1172, 1155, 1047, 980 cm⁻¹; ¹H NMR (DMSO-d₆): δ 10.25 (1H, s, Ar-OH), 9.39 (1H, s, Ar-OH), 8.21 (1H, s, H-2), 7.38 (1H, d, $J=8.8$ Hz, H-5), 7.21 (2H, d, $J=8.3$ Hz, H-2',6'), 6.92 (1H, d, $J=8.8$ Hz, H-6), 6.82 (1H, d, $J=8.3$ Hz, $H=3', 5'$), 3.70 (3H, s, Ar-OCH₃), 3.61 (2H, s, H-9); ¹³C NMR $(CDCl₃+DMSO-d₆)$: δ 176.2, 157.4, 152.1, 149.4, 146.8, 132.5, 130.9, 129.3, 122.7, 117.0, 115.1, 113.7, 113.3, 54.6, 30.0; MS (ESI, negative ion mode): mlz 297 (M-H)⁻. Analysis found: C, 68.41; H, 4.79%. Calcd for $C_{17}H_{14}O_5$: C, 68.45; H, 4.73%.

4.3.8. 5,7-Dihydroxy-3-[(4-hydroxyphenyl)methyl]-4Hchromen-4-one (4h). Pale yellow powder (725 mg, 85%), mp 215–217 °C (lit.^{[19](#page-75-0)} mp 218–219 °C); IR (KBr): 3283, $1667, 1620, 1314, 1282, 1233, 1174, 1053$ cm⁻¹; ¹H NMR (DMSO-d6): d 12.73 (1H, s, Ar-OH), 10.83 (1H, s, Ar-OH), 9.19 (1H, s, Ar-OH), 8.13 (1H, s, H-2), 7.06 (1H, d, $J=$ 8.3 Hz, H-2['],6'), 6.65 (2H, d, $J=8.3$ Hz, H-3',5'), 6.32 (1H, d, $J=2.0$ Hz, H-6), 6.17 (1H, d, $J=2.0$ Hz, H-8), 3.53 (2H, s, H-9); ¹³C NMR (CDCl₃+DMSO- d_6): δ 180.8, 164.0, 161.6, 157.8, 155.6, 153.3, 129.3, 128.5, 122.3, 115.1, 104.4, 98.7, 93.5, 29.3; MS (ESI, negative ion mode): m/z $283 \ (M-H)^{-}$.

4.4. Antioxidant activity

4.4.1. Superoxide free radical scavenging activity. The superoxide free radical scavenging activity was determined by the NBT method. $24,25$ The reaction mixture contained EDTA (6.6 mM), NaCN (3 μ g), riboflavin (2 μ M), NBT (50 μ M), various concentrations of the test drug in ethanol and a phosphate buffer (58 mM, pH 7.8) in a final volume of 3 mL. Optical density was measured at 560 nm. The test tubes were uniformly illuminated with an incandescent lamp for 15 min, after which the optical density was measured again at 560 nm. The percent inhibition of superoxide radical generation was measured by comparing mean absorbance values of the control and those of the test substances. IC_{50} values were obtained from the plot drawn of concentration in μ g versus percentage inhibition and were

converted into μ M. All the tests were run in triplicate and averaged.

4.4.2. DPPH free radical scavenging activity. DPPH radical scavenging activity was measured based on the reduction of methanolic solution of the colored DPPH.^{[26](#page-75-0)} Free radical scavenging ability of the test drug in ethanol added to the methanolic solution of DPPH is inversely proportional to the difference in initial and final absorption of DPPH solution at 516 nm. Drug activity is expressed as the 50% inhibitory concentration (IC_{50}) . The reaction mixture contained 1×10^{-4} mM methanolic solution of DPPH and various concentrations of test drugs. The percentage inhibition was determined by comparing the absorbance values of test and control tubes. IC_{50} values were obtained from the plot, drawn for concentration in microgram versus percentage inhibition.

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Unexpected formation of aryl dialkyl carbinol as a side product from the reaction of methoxyarylaldehydes with Grignard reagents $*$

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Abstract—In the attempted formation of secondary aryl alkyl carbinols from the reaction of methoxyarylaldehydes with Grignard reagents, aryl dialkyl carbinols were formed as unexpected side products. A mechanism for their formation is proposed. Q 2005 Published by Elsevier Ltd.

1. Introduction

Hypercholesterolemia has been accountable for as much as 50% mortality in both developing and developed nations. A number of drugs such as statins, clofibrate and niacins have been developed for improving plasma lipid levels in human beings.^{[1](#page-80-0)} In this regard, some natural phenylpropanoids^{[2](#page-80-0)} $(C_6-C_3$ units) such as α -asarone and elemicin have also evolved as extremely efficient hypolipidemic trans-arylalkenes, although their activity is somewhat marred by genotoxicity of the compounds. As a result, a number of other phenylpropanoids such as arylalkanones and arylalkanols $(C_6-C_3$ units) have been synthesized and have shown promising hypolipidemic activity with reduced toxicity.[2d](#page-80-0) Among these analogues, arylalkanols are important hypolipidemic compounds and are precursors for the synthesis of all above phenylpropanoids. The preparation of arylalkanols^{[3](#page-80-0)} would generally involve reaction of Grignard reagents^{[4](#page-80-0)} with arylaldehydes. Although these Grignard reactions are clean and moderately yielding, formation of various side products^{[5](#page-80-0)} is one problem generally encountered. In most cases, the side products remain unrevealed^{[3a](#page-80-0)} and in others, where they are identified, numerous mechanisms underlying their formation create considerable ambiguity regarding their formation. From an industrial point of view, endeavours towards identification of such side products during the development of chemical

processes for bioactive compounds holds considerable importance, as side product detection is highly crucial for quality approval by relevant agencies. When studying reaction of methoxylated arylaldehydes 2a–2g with alkylmagnesium halides for the formation of aryl alkyl carbinols 3a–3i, we encountered an interesting class of side products, aryl dialkyl carbinol derivatives 1a–1g, along with above expected products 3a–3i ([Scheme 1](#page-77-0)).

2. Results and discussion

In continuation of our ongoing studies on the chemical synthesis of phenylpropanoids^{6} and the associated scale up and yield enhancement for various biological activities, we were interested to prepare intermediate 1-(3,4,5-trimethoxyphenyl)-1-propanol 3a for the synthesis of either 1-(3,4,5-trimethoxyphenyl)-1-propene (commonly known as elemicin) or 1-(3,4,5-trimethoxyphenyl)- 1-propanone. The methodology for the aforementioned synthesis would comprise Grignard reaction of 3,4,5 trimethoxybenzaldehyde 2a with ethylmagnesium bromide in ether to provide 3a, which upon either dehydration or oxidation would provide bioactive^{[2](#page-80-0)} elemicin or phenylpropanone 4a. The first step of Grignard reaction in ether was critically evaluated and this idea was informed by matters emerging in Grignard chemistry wherein a lot of modifications^{$4,5$} have come to the fore. For example, change of the ether solvent in the Grignard process was attempted by replacing this with a mixture of ether and a hydrocarbon of high boiling point such as toluene.[7](#page-80-0) The reaction kinetics and, in turn, yield emanating from such a change of solvent would be

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Keywords: Aryl dialkyl carbinol; Aryl alkyl carbinol; Arylaldehyde; Grignard reaction.

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Scheme 1.

explored. In addition we and others have already reported on formation of methoxylated aryl alkyl carbinols^{[2a,8](#page-80-0)} through Grignard reaction of the corresponding arylaldehydes, wherein the effect of reverse addition of Grignard reagent led to an enhancement in yield of the product carbinols, for example, 3a in 79% yield was observed. In order to further increase the yield of the product 3a, a lot of variants in the reaction (increasing time of addition, changing the amount of the solvents and the amount of the Grignard reagent etc.) were performed. However, success kept eluding us in our endeavour. In due course, we thought that the presence of side product formation in the product mixture was responsible for the comparatively low yield of 3a (79% yield). Unfortunately, we had missed the side product formation during our previous report^{8b} because of performing the Grignard reactions on a small scale (0.012 mol of substrate) and subsequent formation of 1a in minute traces. However, performing the above reactions on a greater preparative scale (0.12 mol of substrate) allowed us to detect the side product 1a in 12% yield. Delving deep into literature studies, one report was found confirming formation of dimer $3c$ as a side product in 30% yield during reaction of arylaldehyde 2b with a large excess of Grignard reagent. This encouraged us to identify the side product, once again, in the present study and attempt to formulate the underlying mechanism for its formation and, subsequently, eliminate

(b) OMe H OMe OMe Me (c) OMe OMe OMe H Me (d) H OMe OMe H Me

(f) OMe H OMe H Me (g) H H OMe H Me (h) OMe H OMe OMe Et (i) OMe H OMe OMe H

 $-OCH₂O- H$ Me

(e) H (R^2+R^3)

production of this side product, so that the yield of the expected product 3a can be improved further.

Surprisingly, the side product formed during reaction of 3,4,5 trimethoxybenzaldehyde 2a with ethylmagnesium bromide was not the expected dimer^{[3c](#page-80-0)} (as reported in the case of $2,4,5$ trimethoxybenzaldehyde 2b) but instead 3-(3,4,5-trimethoxyphenyl)-3-pentanol 1a, a tertiary carbinol. Purification of the crude product by column chromatography provided 3a in 79% and $1a$ in 12% yield. The ¹H and ¹³C NMR spectra of 3a matched well with the reported values.^{[8b](#page-80-0)} The ¹H NMR spectrum of 1a indicated a 6H triplet at δ 0.73 and also a quartet of 4H at 1.76, which indicated the presence of 2 equiv $-CH_2-CH_3$ groups. The presence of two ethyl groups was further confirmed by the DEPT 13 C NMR spectrum of 1a. The DEPT spectrum in d_6 -DMSO confirmed the point of attachment of the additional $-CH_2-CH_3$ group at C-1 as the signal at 77.2 disappeared due to its quaternary nature. HRMS of 1a also supported the proposed structure of the aryl dialkyl carbinol.

The formation of the unexpected product 1a was unequivocally confirmed by its synthesis starting from 3a. Oxidation of $3a$ with $DDQ⁹$ $DDQ⁹$ $DDQ⁹$ afforded $4a$ in 89% yield, which upon treatment with ethylmagnesium bromide provided 1a in 78% yield (Scheme 2), whose TLC, mixed TLC, co-mp and NMR spectra were identical to side product 1a discussed above.

Scheme 3.

Table 1. Grignard reaction of arylaldehydes (2a–2i) into aryl alkyl carbinols (3a–3i) and unexpected side product aryl dialkyl carbinols (1a–1g)

Run	Arylaldehyde (2)	Aryl alkyl carbinol (3)	Aryl dialkyl carbinols (1)
\rm{a}	CHO	QH	QН
	H_3CO	H_3CO	H_3CO
	H_3 CO	H_3CO	H_3CO
	$\overline{O}CH_3$	$\overline{O}CH_{3}$ _{79%}	OCH_3 12%
$\mathbf b$	QCH ₃ CHO H_3 CO $\overline{O}CH_3$	QCH ₃ QH H_3 CO $\frac{1}{2}$ CH ₃ $\frac{79\%}{2}$	QCH ₃ OH H_3CO 12%, \overline{OCH}_3
$\mathbf c$	QCH ₃	QCH ₃ QH	$OCH3$ OH
	CHO	H_3CO	H_3CO
	H_3CO	H_3CO	7%
	H_3CO	84%	H_3 CO
$\mathrm{d}% \left\ \mathcal{H}\right\ _{A}$	H_3CO CHO H_3CO	QH H_3CO 81% H_3CO	QН H_3CO H_3CO 8%
$\rm e$	CHO	QН 74% Ō , 88%	OH 7%
$\mathbf f$	QCH ₃	QCH ₃ QH	QCH ₃ QH
	CHO	79%	H_3CO
	H_3CO	H_3CO	8%
g	CHO H_3CO	QН 78% H_3CO	QH H_3CO 8%
h	QCH ₃	QCH ₃ QH	QCH ₃ OH
	CHO	85%	H_3 CO
	H_3 CO	H_3 CO	0%
	QCH ₃	ϕ CH ₃	\overline{OCH}_3
$\rm i$	QCH ₃	QCH ₃ QH	QCH ₃ OH
	CHO	87%	H_3 CO
	H_3 CO	H_3 CO	0%
	\overline{OCH}_3	$\overline{O}CH3$	QCH ₃

The mechanism for formation of dialkyl carbinol 1a is believed to proceed through an internal Canninzaro-type reaction^{[3c](#page-80-0)} of some of the secondary alcohol **3a** to afford the corresponding oxidized product 4a [\(Scheme 3\)](#page-78-0).

After successful isolation of the side product 1a, the Grignard reaction was performed with 2,4,5-trimethoxybenzaldehyde 2b and, once again, tertiary carbinol (1b) was obtained as a side product in 12% yield but not the dimer as previously isolated by Francisco et al. 3^b The plausible reason for the absence of any dimeric side product in our case may be due to the reverse addition of Grignard reagent to 2b, in comparison to the reported method.^{[3b](#page-80-0)} In order to check the generality of the above reaction, some other substituted arylaldehydes 2c–2g were also studied and the formation of the tertiary alcohols 1c–1g was detected along with the major products 3c–3i [\(Table 1](#page-78-0)). In contrast, no aryl dialkyl carbinols 1h–1i were detected in the case of Grignard reaction with both propylmagnesium bromide and methylmagnesium bromide, and arylaldehyde 2b. However, the normal aryl alkyl carbinols 3h–3i were both formed in these cases in better yields than the other reactions described. This finding further strengthens our hypothesis of formation of the side product at the cost of the normal expected product. Further studies to eliminate the side product 1a–1g in order to enhance the yield of carbinols 3a–3g are in progress.

3. Conclusion

In conclusion, we have discovered the formation of aryl dialkyl carbinols 1a–1g from the reaction of methoxyarylaldehydes 2a–2g with alkylmagnesium halides during the formation of aryl alkyl carbinols 3a–3i. The formation of tertiary alcohols is reported from acyl, carboxylic acid, ketone and ester derivatives, but their direct formation from aldehydes is a new finding. Overall, identification of such side products during development of chemical processes for the synthesis of bioactive compounds holds much importance.

4. Experimental

4.1. General

Melting points were determined with a Mettler FP80 micromelting point apparatus and are uncorrected. Column chromatography was performed on silica gel (60–120 mesh size). ¹H (300 MHz) NMR spectra were recorded in CDCl₃ and d_6 -DMSO on a Bruker Avance-300 spectrometer. HRMS were determined using a Micromass Q-TOF Ultima spectrometer.

4.2. Representative experimental for the synthesis of 3a–3i and 1a–1g

Alkylmagnesium bromide (prepared from 30.5 mL of alkyl bromide and 9.91 g Mg in 70 mL of ether) in toluene (40 mL) was added dropwise in 15–20 min to a cooled mixture of methoxy arylaldehyde (2a–2g, 0.102 mol) in ether (80 mL) and toluene (125 mL) and finally the mixture was stirred for 12–14 h at room temperature under a nitrogen atmosphere. The cooled mixture was poured into a saturated solution of ammonium chloride $(2 \times 15 \text{ mL})$ and stirred for 20 min. The mixture was extracted with ethyl acetate $(3 \times 50 \text{ mL})$ and the combined extracts were washed with brine (2×15 mL), dried over Na₂SO₄ and filtered. The residue obtained upon evaporation was chromatographed by neutral alumina column using a hexane–ethyl acetate mixture, with increasing proportions of ethyl acetate up to 25%, to afford pure secondary alcohols 3a–3i and tertiary alcohols 1a–1g. Spectral data for aryl alkyl carbinols $(3a-3i)$ was found to match with the reported values.^{[8b](#page-80-0)}

4.2.1. Compound 1a. White solid, 3.1 g (12%); mp 60– 62 °C; $v_{\text{max}}(\text{CHCl}_3)$ 3571 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): d 6.52 (2H, s, H-2 and H-6), 3.78 (9H, s, 3,4,5-OCH₃), 1.76 (4H, q, $J = 6.85$ Hz, H-2^f and H-2^f), 0.73 (6H, t, $J=6.85$ Hz, H-3['] and H-3[']); ¹³C NMR (CDCl₃, 75.4 MHz): d 153.05 (C-3 and C-5), 142.2 (C-4), 136.5 (C-1), 103.2 (C-2 and C-6), 77.2 (C-1'), 61.1 (4-OCH₃), 56.4 (3, 5-OCH₃), 35.3 (C-2['] and C-2[']), 8.2 (C-3['] and C-3[']); HRMS (M+Na) m/z : 277.3176 (Calcd for C₁₄H₂₂O₄Na: 277.3160).

4.2.2. Compound 1b. White solid, 3.1 g (12%); mp 107–110 °C; $\nu_{\text{max}}(\text{CHCl}_3)$ 3568 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): d 7.02 (1H, s, H-6), 6.38 (1H, s, H-3), 3.83 (6H, s, 4 and 5-OCH₃), 3.79 (3H, s, 2-OCH₃), 2.34 (4H, q, $J=$ 7.3 Hz, H-2' and H-2'), 1.12 (6H, t, $J=7.3$ Hz, H-3['] and H-3'); ¹³C NMR (CDCl₃, 75.4 MHz): δ 160.0 (C-2), 156.5 (C-4), 141.8 (C-5), 111.1 (C-1), 110.9 (C-6), 100.6 (C-3), 78.0 (C-1'), 56.7 (4, 5-OCH₃), 56.1 (C-2-OCH₃), 31.3 (C-2['] and C-2'), 8.3 (C-3' and C-3'); HRMS (negative) m/z : 253.3201 (Calcd for $C_{14}H_{22}O_{4}$: 253.3182).

4.2.3. Compound 1c. Colourless liquid, 1.8 g (7%); $\nu_{\text{max}}(\text{CHCl}_3)$ 3575 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): δ 6.80 (1H, d, $J=8.9$ Hz, H-6), 6.48 (1H, d, $J=8.9$ Hz, H-2), 3.81 (3H, s, 3-OCH3), 3.70 (6H, s, 2 and 4-OCH3), 1.87-1.75 (2H, m, H-2'), 1.69-1.57 (2H, m, H-2'), 0.63 (6H, t, $J=7.3$ Hz, H-3^t and H-3^t); ¹³C NMR (CDCl₃, 75.4 MHz): d 152.3 (C-5), 151.8 (C-4), 142.1 (C-3), 128.9 $(C-6)$, 122.0 $(C-1)$, 106.5 $(C-2)$, 78.0 $(C-1')$, 60.6 (3-OCH₃), 60.4 (2-OCH₃), 55.7 (4-OCH₃), 33.9 (C-2['] and C-2[']), 8.2 $(C-3)$ and $C-3$ [']); HRMS $(M + Na)$ m/z: 277.3158 (Calcd for $C_{14}H_{22}O_4$ Na: 277.3160).

4.2.4. Compound 1d. Colourless liquid, 1.8 g (8%); $v_{\text{max}}(\text{CHCl}_3)$ 3572 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): δ 6.82 (1H, d, $J=8.1$ Hz, H-6), 6.74 (1H, d, $J=8.1$ Hz, H-5), 6.24 (1H, s, H-2), 3.82 (6H, s, 3 and 4-OCH3), 1.81–1.78 (4H, m, H-2' and H-2'), 0.92 (6H, t, $J=7.3$ Hz, H-3' and H-3'); ¹³C NMR (CDCl₃, 75.4 MHz): δ 149.0 (C-3), 148.1 (C-4), 131.1 (C-1), 130.6 (C-6), 123.8 (C-5), 118.6 (C-2), 77.2 (C-1'), 55.9 (4-OCH₃), 55.7 (3-OCH₃), 29.7 (C-2' and C-2'), 13.2 (C-3' and C-3'); HRMS (posative) m/z : 225.3012 (Calcd for $C_{13}H_{20}O_3$: 225.3079).

4.2.5. Compound 1e. Colourless liquid, 1.5 g (7%); $\nu_{\text{max}}(\text{CHCl}_3)$ 3567 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): δ

6.83 (1H, s, H-6), 6.76 (2H, m, H-2 and H-5), 5.91 (2H, s, $-OCH_2O$ –), 1.79–1.73 (4H, m, H-2' and H-2'), 0.89 (6H, t, $J=6.9$ Hz, H-3' and H-3'); ¹³C NMR (CDCl₃, 75.4 MHz): δ 149.0 (C-3), 148.2 (C-4), 134.5 (C-1), 121.6 (C-6), 112.3 $(C-5)$, 105.1 $(C-2)$, 98.5 $(-OCH₂O₋)$, 77.1 $(C-1')$, 30.7 $(C-2')$ and $C-2'$), 11.4 $(C-3'$ and $C-3'$); HRMS (negative) m/z : 207.2419 (Calcd for $C_{12}H_{16}O_3$: 207.2493).

4.2.6. Compound 1f. Colourless liquid, 1.8 g (8%); v_{max} (CHCl₃) 3571 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): δ 7.20 $(1H, d, J=8.5 Hz, H=6)$, 6.5 (1H, d, $J=8.5 Hz, H=5)$, 6.39 $(1H, s, H-3), 3.69$ (6H, s, 2 and 4 – OCH₃), 1.71–1.69 (4H, m, H-2' and H-2'), 0.92 (6H, t, $J=7.3$ Hz, H-3' and H-3'); ¹³C NMR (CDCl₃, 75.4 MHz): δ 159.7 (C-2), 158.8 (C-4), 129.4 (C-6), 117.2 (C-1), 102.2 (C-5), 99.1 (C-3), 72.6 $(C-1'), 55.8 (2- OCH₃), 55.2 (4- OCH₃), 33.1 (C-2' and C-2'),$ 8.1 (C-3['] and C-3[']); HRMS (positive) m/z: 225.3012 (Calcd for $C_{13}H_{20}O_3$: 225.3079).

4.2.7. Compound 1g. Colourless liquid, 1.6 g (8%); $v_{\text{max}}(\text{CHCl}_3)$ 3571 cm⁻¹; ¹H NMR (CDCl₃, 300 MHz): δ 7.21 (2H, d, $J=8.5$ Hz, H-6 and H-2), 6.79 (2H, d, $J=$ 8.5 Hz, H-5 and H-3), 3.72 (3H, s, $4'$ -OCH₃), 1.74 (4H, m, H-2' and H-2'), 0.69 (6H, t, $J=7.3$ Hz, H-3['] and H-3'); ¹³C NMR (CDCl₃, 75.4 MHz): δ 158.0 (C-4), 137.9 (C-1), 126.6 $(C-6$ and $C-2)$, 113.3 $(C-5$ and 3), 77.1 $(C-1')$, 55.1 $(4\text{-}OCH_3)$, 34.8 $(C-2'$ and $C-2'$), 7.8 $(C-3'$ and $C-3'$); HRMS (M+Na) m/z : 217.2595 (Calcd for C₁₂H₁₈O₂Na: 217.2634).

4.3. Synthesis of 1-(3,4,5-trimethoxy)phenyl-1-propanone (4a) and 1a from 3a

A mixture of aryl alkyl carbinol (3a) (2.4 mmol) and DDQ (4.8 mmol) in wet dioxane (30 mL) was stirred at room temperature for 14 h. The precipitated $DDQH₂$ was filtered and the red coloured filtrate was evaporated and subsequently chromatographed on silica gel (hexane/ethyl acetate 7:3) to provide phenylpropanone 4a in 89% whose spectral data was found matching with the reported values. Further treatment of ketone 4a with excess of ethylmagnesium bromide in ether and toluene provided 1a in 78% yield.

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Supplementary data

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Phase transfer catalyzed aziridination of a-bromo-2-cyclopenten-1-one

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Abstract—An efficient and highly selective synthesis of bicyclic-a-keto aziridines from 2-bromo-2-cyclopentenone and aliphatic primary amines mediated by phase transfer catalysts (PTCs) in water at room temperature is demonstrated. Bicyclic- α -keto-aziridines are highly strained and reactive compounds that can be used in the synthesis of biologically active compounds. Therefore, the present strategy with its mild reaction conditions opens up a new entry to the synthesis of unusual aziridines using inexpensive reagents. Q 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Three-membered-heterocycles such as epoxides and aziridines are versatile intermediates in organic synthesis because they can be converted into a variety of multi-functional compounds.^{[1,2](#page-84-0)} The ability of aziridines to undergo highly regio- and stereoselective ring-opening reaction renders them useful as precursors for chiral substrates, auxiliaries, reagents and ligands in stereoselective synthesis for a variety of N -containing compounds.^{[3,4](#page-84-0)} The nucleophilic ring-opening of N-substituted aziridines in the presence of Lewis acids regioselectively affords substituted α -functionalized β -amino acid or β -functionalized α -amino acid precursors, depending on the nature of the nucleophile, the Lewis acid and the three-membered ring substituents. Through ring expansion, aziridines could also give amido derivatives and oxazolines, as a protected form of hydroxyl amino compounds.^{[5–7](#page-84-0)} In addition. b-lactam antibiotics, pyrrolidine alkaloids, polymers and chiral amines as chiral auxiliaries for asymmetric alkylation and aldol transformations could be obtained from aziridines.[8,9](#page-84-0) Enantiopure aziridines are also currently of interest as enzyme substrates and enzyme inhibitors.^{[9](#page-84-0)} Consequently, novel types of peptidic cysteine protease inhibitors containing aziridine-2,3-dicarboxylic acid have been designed and recently synthesized.^{[10](#page-84-0)}

The development of efficient synthetic routes to aziridines is therefore a worthy target for the synthetic organic chemist.^{[11,12](#page-84-0)} It is essential that efficient methods exist for the facile synthesis of a range of structurally diverse aziridines, with the added requirement that any available methods should also allow stereoselective aziridine formation.[1](#page-84-0) Like their epoxide counterparts, aziridines can be prepared by a number of methods most of which start from easily available enantiomerically pure compounds. Olefins, imines and β -amino alcohols are the most common and attractive precursors of aziridine derivatives due to low cost, wide availability and their ability to undergo direct $[2+1]$ aziridination reactions[.12a](#page-85-0)

Although the stereoselective synthesis of activated aziridines from olefins, alcohols and imines has been widely investigated, $13-15$ to the best of our knowledge, studies on non-activated aziridines (N-alkyl or aryl) are very rare.^{[3,16](#page-84-0)} Aziridination of 2-iodo-a, b-unsaturated ketones using $Cs₂CO₃$ and xylene in the presence of 1,10-phenanthroline refluxing at 90 \degree C was reported by Barros and co-workers.^{[3](#page-84-0)} We have, however, found that bicyclic-a-keto-aziridines are unstable and decompose to form non-homogenous mixtures within few days even when kept in a refrigerator. This shows that their synthesis requires mild reaction conditions in order to make it possible to isolate reasonable yields.

Nowadays, the demand for environmentally benign organic chemical transformations and organizating research with the aim of keeping pollution effects to a minimum, together with a reduction of energy and raw materials consumption has increased.^{[17,18](#page-85-0)} In this respect, the procedure developed by Barros et al.^{[3](#page-84-0)} suffers from several disadvantages such as

Keywords: Aziridination; Phase transfer catalysis; Primary aliphatic amines; Conjugate addition-imitated ring closure reaction; Bicyclic-aketo-aziridines.

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long reaction time, use of toxic and organic solvent, difficulty of recovery of high boiling solvent relative to the stability of aziridines, high reaction temperature and use of additives. These indicate that a better method for this transformation is desired. We have developed a practical, highly efficient and simple aziridinaton protocol of a-bromo-2-cyclopenetenone 2. We have assessed different reaction conditions to apply tandem conjugate additioninitiated ring closure (CAIRC) reactions and in the present paper we wish to report the synthesis of N-substituted bicyclic-a-keto-aziridines from a series of primary amines applying various PTCs particularly using tetrabutylammonium bromide, TBAB in H_2O .

2. Results and discussion

One of the long standing goals of our research has been the development of a mild and general protocol for the selective functionalization of α -donor substituted 2-cyclopentenones 2 with different nucleophiles. Our recent findings have led us to pose several fundamental questions that relate to the stereoselective synthesis of N, S and O-containing heterocycles through conjugate addition-initiated ring closure reactions (CAIRC). Specifically, we are interested in the construction of aziridines from amines since N-substituted aziridines are important building blocks for the synthesis of α and β -amino acids. It occurred to us that 2-bromo-2cyclopentenone 2 could be a suitable substrate for the synthesis of bicyclic a-keto-aziridines.

Initially we chose a model reaction: the addition of benzylamine 1a to 2 and we investigated different molar ratio of reactants. We also scanned several non-polar and polar solvents such as CH_2Cl_2 , CH_3CN , THF, Et_2O , $EtOAc$, toluene, C_6H_6 , CH₃OH, acetone and CHCl₃, as well as H₂O. It was noteworthy that the reaction proceeded smoothly only in $H₂O$. In all the cases, the presence of PTC was found to be essential to promote these reactions. Therefore, we further explored the catalytic activities of various quaternary ammonium salts in the presence of water in tandem conjugate addition-initiated ring closure reaction to give aziridines and the results are summarized in Table 1. The experiments reported in Table 1 were all carried out in water and yields were not optimized. Most of the reactions gave similar results using various PTCs. Especially, the aziridination reactions mediated by Bu_4NBr , Bu_4NI , Hex_4NBr and benzyldimethyl-2-hydroxyethyl ammonium chloride (entries 1, 3, 6 and 8, respectively) proceeded in a similar manner with comparable yields. Bu4NF showed slow

Table 1. Screening of some phase transfer catalysts on aziridination of 2 with the model amine $1a$ in the presence of H_2O at room temperature

Entry	Phase transfer catalysts	Yield $(\%)$
	Tetrabutylammonium bromide	91
\overline{c}	Tetrabutylammonium fluoride	54
3	Tetrabutylammonium iodide	88
$\overline{4}$	Tetrabutylammonium tetrafluoroborate	79
5	Tetrabutylammonium hydrogensulphate	82
6	Tetrahexylammonium bromide	87
	Benzyltriaethylammonium chloride	74
8	Benzyldimethyl 2-hydroxyethylammonium chloride	89

conversion in which most unreacted materials were recovered together with significant amount of amination products. However, for convenience and simplicity, we chose TBAB in $H₂O$ to perform the aziridination reaction of 2-bromo-2-cyclopentenone 2 with other amines.

After choosing water as solvent and TBAB as catalyst, we next observed the reaction between 2 and 1a at room temperature. The reactions were fast giving high yields of the desired products after a reasonable time of reaction. When the reactions were run in water using TBAB as a catalyst, we found that the stoichiometry of the reactants was critical. The use of excess amounts or even equimolar amount of amines leads to substantial amounts of byproducts through transamination or alkylation or bisaddition. This is probably due to HBr liberated during the reaction that protonates the aziridine formed, which is subsequently ring opened by the excess amine (Scheme 1). Therefore, the relief of ring-strain is a factor that provides a driving force for ring-opening after protonation. Primary amines can also be protonated by HBr to give self-alkylated tri- and diamines. For instance, when excess 1c was reacted with 2, $(C_5H_5O)_3N$ was obtained as a major byproduct, which was identified by GC–MS.

Scheme 1.

When a slight excess of acceptor (1:1.25 mmol) and 10 mmol % TBAB was used, the side reactions was totally suppresses and no byproducts were obtained. This showed that the course of the reaction was highly dependent on the amount of reactants used. Slight excess of enone provided a good compromise between conversion, yield and suppression of side products so that aziridines can be obtained in 90–98% isolated yields in reactions carried out at room temperature.

To determine the scope and generality of the protocol with respect to amines variation, a series of aliphatic primary amines 1a–h (phenethyl, benzyl, furayl, cyclohexyl, allyl, butyl, tert-butyl and propyl amines, respectively) were reacted with 2 under the above-mentioned conditions and the results are summarized in [Table 2](#page-83-0). As shown in [Table 2](#page-83-0), we observed very good to excellent reactivity with all the tested amines. Particularly, the reaction of t -BuNH₂ was very fast and the reaction was completed within 1 h. The use of aromatic amines such as aniline, 2, 3-dimethylaniline and 2, 6-dimethylaniline under the same reaction conditions did not produce any aziridines nor even after 3 days of stirring. This can be explained by the lower solubility and the less nucleophilc nature of the aromatic amines compared to aliphatic amines towards the first conjugate addition. The effect of solubility in water is also clearly seen among the reactivity of aliphatic amines. Aliphatic amines containing aromatic groups in their chain were found to be relatively less reactive than others ([Table 2,](#page-83-0) entries 1, 2 and 3). This

Scheme 2.

Table 2. TBAB catalyzed aziridination of 2 with amines $1a-h$ in H₂O at room temperature

Entry	Amine	Time (h)	Product	Yield (%)
$\,1$	1a	5	Ö 3a	98
$\boldsymbol{2}$	1 _b	5	3 _b	93
$\sqrt{3}$	1c	5	٠N 3c	91
$\overline{\mathcal{L}}$	1 _d	$\mathfrak z$	3d	90
5	1e	$\overline{\mathbf{3}}$	N. 3e	96
$\boldsymbol{6}$	1f	$\mathfrak z$	3f	95
$\boldsymbol{7}$	$1\mathrm{g}$	$\,1$		94
$\,$ 8 $\,$	1 _h	3	3g 3 _h	97

indicates that the quaternary ammonium salts are suitable catalysts to active aliphatic amines and not aromatic amines. This observed selectivity could be useful to differentiate the two types of amines for synthetic applications (Scheme 2).

Generally, this procedure is used to introduce aliphatic amines having different functionality such as allyl and furfuryl groups. Particularly, allyl containing aziridines are useful and versatile synthetic intermediates since this group can be derivatized into interesting functionalities.

3. Conclusion

In this paper, we show that highly reactive bicyclic α -ketoaziridines can be synthesized from α -bromo-enones via a simple 'one pot' procedure. This is the first strategy where aziridines have been synthesized from 2-bromo-2-cyclopentenone and amines mediated by PTCs in water. We believe that this method offers advantages for the development of a direct diastreoselective synthesis of substituted aziridines in high yields under very mild and simple conditions. Further studies will focus on the development of asymmetric aziridination reaction of amines and on the extension of the reaction scope to other amines.

4. Experimental

4.1. General methods

All 1 H and 13 C NMR spectra were recorded on a JEOI JNM-EX 400 FT NMR system using CDCl₃ as a solvent. Chemical shifts are given in ppm and J-values in Hz. Analytical TLC were carried out on precoated (0.25 mm) Merck silica gel F-254 plates. Flash chromatography was carried out using Merck's Kiegelgel 60 (230–400mesh). GLC analyses were performed on a varian 3300 chromatograph equipped with split injector, FID detector and a varian 4400 integrator. IR spectra were recorded on a FT-IR spectrometer and are reported as wave number. GC–MS spectra were registered on a Hewlett 5890 Packard series II CP Sil 5 CB column (25 m) followed by VG Quattro mass spectrometer. Finnigan-MAT-95XL mass spectrometer was used to obtain HREIMS data and the spectra were obtained at 250° C and 70 eV . All reagents and solvents except 2-bromo-cyclopentenone 2 were obtained from commercial sources and used as received without further purification. Compound 2 was prepared according to literature procedure.[19](#page-85-0)

4.2. General procedure for the reaction of 2-bromo-2 cyclopenten-2-one 2 with various amines (1a–h)

A mixture of the bromo-ketone 2 (1.25 mmol), TBAB (10 mmol %) and aliphatic primary amines (1.0 mmol) in $H₂O$ (5 mL) was stirred at room temperature for the specified time until the amine was consumed (see Table 2). The reaction mixture was extracted with $Et₂O$ $(3 \times 5 \text{ mL})$. The combined organic layers were dried over $Na₂SO₄$ and the solvent was removed under reduced pressure. The expected aziridines (3a–h) were obtained in quantitative yields after flash chromatography on silica gel (pentane/diethyl ether, 4:1). Each aziridine was characterized by the analysis of HREIMS, DEPT, ¹H and $13C$ NMR data except 3a and 3e that showed very short life spans in the HREIMS.

4.2.1. 6-Benzyl-6-aza-bicyclo[3.1.0]hexan-2-one (3a). (0.181 g, 98%) as a brown oil, R_f 0.46 (3:7 Et₂O/pentane); IR (neat, NaCl plates, ν_{max} , cm⁻¹): 3028, 2930, 1738, 1601, 1451, 1351; ¹H NMR (δ): δ 1.95–2.10 (m, 2H), 2.24–2.35 $(m, 2H)$, 2.40 (d, J=4.0 Hz, 1H), 2.80 (t, J=3.6 Hz, 1H), 3.40 (d, $J=13.6$ Hz, 1H), 3.80 (d, $J=13.6$ Hz, 1H), 7.27–7.34 (m, 1H), 7.35–7.42 (m, 4H); ¹³C NMR: δ 212.1 (CO), 139.7 (C), 129.2 (CH), 128.5 (CH), 127.3 (CH), 60.2 $(CH₂Ph)$, 47.6 (CH), 46.5 (CH), 32.6 (CH₂), 24.2 (CH₂); MS (EI): 187% (M⁺, 35), 131 (15), 92 (83), 91 (100), 77 (16), 65 (89), 51 (21).

4.2.2. 6-(Phenethyl)-6-aza-bicyclo[3.1.0]hexan-2-one (3b). (0.187 g, 93%) as brown oil, R_f (0.41 Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2929, 1712, 1421, 1360, 1220; ¹H NMR (δ): δ 1.82–2.20 (m, 2H), 2.06 (d, J = 4.0 Hz, 1H), 2.13–2.21 (m, 1H), 2.25–2.35 (m, 1H), 2.46 (t, $J=$ 3.6 Hz, 1H), 2.52–2.60 and 2.63–2.71 (m, $-CH₂Ph$), 2.92 $(t, J=7.6 \text{ Hz}, 2\text{H}), 7.18-7.26 \text{ (m, 2H)}, 7.28-7.39 \text{ (m, 3H)};$ 13 C NMR: δ 212.0 (–CO), 139.5 (C), 129.0 (CH), 128.5 (CH), 126.2 (CH), 60.2 (CH), 47.0 (CH), 46.4 (CH₂), 36.5 (CH_2) , 32.6 (CH_2) , 24.2 (CH_2) ; HRMS: found for C₁₃H₁₅NO: 201.1152, calcd: 201.1154.

4.2.3. 6-Furan-2-ylmethyl-6-aza-bicyclo[3.1.0]hexan-2-one (3c). (0.162 g, 91%) as a brown oil, R_f 0.79 (3:7 Et₂O/ pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 3116, 2932, 1739, 1590, 1503, 1446, 1405, 1344; ¹H NMR (δ): δ $1.96-2.05$ (m, 2H), $2.22-2.36$ (m, 2H), 2.35 (d, $J=3.6$ Hz, 1H), 2.82 (t, $J=3.6$ Hz, 1H), 3.42 (d, $J=14.2$ Hz, 1H), 3.63 (d, $J=14.2$ Hz, 1H), 6.13 (d, $J=3.0$ Hz, 1H), 6.22 (dd, $J=3.0$, 1.0 Hz, 1H), 7.17 (d, $J=1.0$ Hz, 1H); ¹³C NMR: δ 212.0, 152.2 (C), 142.5 (CH), 110.5 (CH), 108.3 (CH), 54.1 (-NCH₂), 46.8 (CH), 45.3 (CH), 33.1 (CH₂), 24.2 (CH₂); HRMS: found for $C_{10}H_{11}NO_2$: 177.0796, calcd: 77.0790.

4.2.4. 6-Cyclohexyl-6-aza-bicyclo[3.1.0]hexan-2-one (3d). (0.161 g, 90%) as a yellow oil, R_f 0.72 (1:4) Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2926, 2853, 1738, 1449, 1353, 1165, 1112; ^TH NMR (δ): δ 1.12–1.26 (m, 2H), 1.28–1.48 (m, 4H), 1.59–1.68 (m, 4H), $1.75-1.89$ (m, 2H), $1.94-2.04$ (m, 1H), 2.10 (d, $J=$ 3.6 Hz, 1H), 2.14–2.36 (m, 2H), 2.62 (t, $J=3.6$ Hz, 1H); ¹³C NMR: δ 212.3, 66.5 (–NCH), 46.2 (CH), 45.4 (CH), 33.5 (CH₂), 33.0 (CH₂), 24.9 (CH₂), 24.7 (CH₂), 24.5 (CH₂); HRMS: found for $C_{11}H_{17}NO$: 179.1314, calcd: 179.1310.

4.2.5. 6-Allyl-6-aza-bicyclo[3.1.0]hexan-2-one (3e). (0.131 g, 96%) as a yellow oil, R_f 0.62 (1:9 Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2929, 1738, 1642, 1448;
¹H NMP (A): λ 1.02, 2.04 (m, 2H), 2.13 (d, $I = 4.0$ H_z, 1H) ¹H NMR (δ): δ 1.92–2.04 (m, 2H), 2.13 (d, J = 4.0 Hz, 1H), 2.20–2.36 (m, 2H), 2.63 (t, $J=3.6$ Hz, 1H), 2.99 (d, $J=5.6$ Hz, 2H), 5.17 (dd, $J=9.2$, 1.2 Hz, 1H), 5.26 (dd, $J=16.0$, 1.6 Hz, 1H), 5.70–5.95 (m, $CH=CH₂$); ¹³C NMR: δ 212.0, 134.4 (CH=CH₂), 117 (CH=CH₂), 60.6 (-NCH₂), 47.0 (CH), 46.2 (CH), 33.3 (CH₂), 24.2 $(CH₂)$; MS: m/z % 137 (M⁺, 25), 136 (M⁺ -H, 6), 71 (11), 57 (12), 41 (24).

4.2.6. 6-Butyl-6-aza-bicyclo[3.1.0]hexan-2-one (3f). $(0.145 \text{ g}, 95\%)$ as a yellow oil, R_f 0.52 (1:9 Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2930, 2867, 1739, 1615, 1456, 1407, 1353; ¹H NMR (δ): δ 0.90 (t, J = 14.8 Hz, 3H) 1.32–1.41 (m, 2H), 1.50–1.60 (m, 2H), 1.88–1.97 (m, 2H), 2.02 (d, $J=4.0$ Hz, 1H), 2.14–2.26 (m, 2H), 2.31 (t, $J=$ 6.8 Hz, 2H), 2.54 (t, $J=3.6$ Hz, 1H); ¹³C NMR: δ 212.2, 58.5 (–NCH₂), 47.2 (CH), 46.1 (CH), 33.2 (CH₂), 32.5 (CH_2) , 24.1 (CH₂), 20.5 (CH₂), 14.3 (CH₃); HRMS: found for C9H15NO: 153.1155, calcd: 153.1154.

4.2.7. 6-tert-Butyl-6-aza-bicyclo[3.1.0]hexan-2-one (3g). $(0.143 \text{ g}, 94\%)$ as a yellow oil, R_f 0.55 (1:9 Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2968, 1737, 1612, 1362,

1213, 1091; ¹H NMR (δ): δ 1.03 (s, 9H), 1.92-2.14 (m, 2H), 2.27 (d, $J=4.0$ Hz, 1H), 2.33–2.42 (m, 2H), 2.78 (t, $J=$ 3.6 Hz, 1H); 13C NMR: d 214.5, 54.2 (C), 41.5 (CH), 39.4 (CH), 35.7 (CH₂), 27.8 (CH₃), 24.5 (CH₂); GC–MS: $m/z\%$ 154 (M⁺ + H, 2), 153 (M⁺, 5), 97 (100), 68 (34), 57 (34), 41 (30).

4.2.8. 6-Propyl-6-aza-bicyclo[3.1.0]hexan-2-one (3h). (0.135 g, 97%) as a yellow oil, R_f 0.69 (1:9 Et₂O/pentane); IR (neat, NaCl plates, v_{max} , cm⁻¹): 2958, 1740, 1680, 1633, 1460, 1352, 1190; ¹H NMR (δ): δ 0.94 (t, J = 14.8 Hz, 3H), $1.54-1.66$ (m, 2H), $1.90-1.97$ (m, 2H), 2.03 (d, $J=3.6$ Hz, 1H), 2.10–2.24 (m, 2H), 2.30 (t, $J=11.2$ Hz, 1H), 2.55 (t, $J=3.6$ Hz, 2H); ¹³C NMR: δ 212.2, 60.5 (–NCH₂), 37.2 (CH), 36.2 (CH), 33.0 (CH₂), 24.1 (CH₂), 23.3 (CH₂), 12.4 (CH₃); HRMS: found for $C_8H_{13}NO: 139.0998$, calcd: 139.0997.

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A microwave-enhanced, solventless Mannich condensation of terminal alkynes and secondary amines with para-formaldehyde on cuprous iodide doped alumina

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Abstract—A microwave-enhanced, solventless Mannich condensation of terminal alkynes and secondary amines with *para*-formaldehyde on cuprous iodide doped alumina has been developed. b-Aminoalkynes are generated in good yields. The reaction can be extended to include a cyclization, which affords 2-substituted benzo $[b]$ furans. The chemoselectivity of the reaction indicates that terminal alkynes are much more reactive than enolizable ketones under the reaction conditions.

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1. Introduction

The Mannich reaction is a classic example of a three-component condensation reaction.^{[1](#page-96-0)} In general, formaldehyde (or para-formaldehyde), an amine, and an 'active-hydrogen' component such as an enolizable ketone or terminal alkyne are allowed to react to afford the corresponding β -aminoketone or β -aminoalkyne. The latter Mannich adduct contains at least two potential sites for further modification: the amine and the alkyne. 2 2 2 In addition, b-aminoalkynes and their derivatives have a wide range of applications including use as pharmaceutical intermediates 3 and as general synthetic building blocks.^{[4](#page-96-0)} Moreover, the alkyne moieties may be functionalized in various ways.

The traditional Mannich method for synthesizing β -aminoalkynes often requires drastic reaction conditions and generally is run in dioxane, a toxic solvent. The organic solvent and the metal catalyst can be difficult to handle and often difficult to dispose of safely.

We have found alumina to be a particularly useful reagent in organic synthesis because it can be modified in a variety of ways that enhance its reactivity. It also obviates a number of environmental problems.^{[5](#page-96-0)} For example, using a commercially available alumina/potassium fluoride mixture to which palladium powder was added, we were able to carry out Suzuki and Sonogashira coupling reactions on a wide variety of aromatic moieties without the use of solvents.^{[6](#page-96-0)}

Microwave irradiation of organic reactions has gained in popularity in recent years since it was found to accelerate a wide variety of transformations.^{[7](#page-96-0)} Early experiments utilized solvents with high dielectric constants, which permitted rapid heating of the reaction solution. In recent years, a number of reports have appeared in which reactants are coated on to surfaces, which themselves absorb little or no microwave energy; in these instances, the reactive species absorb the microwave energy but the temperature of the reaction mixture tends to rise only modestly. This results in relatively large energy saving as well as making it possible to carry out reactions in relatively simple glassware such as open beakers and flasks.^{[8](#page-96-0)}

We now wish to report the details of a microwave-enhanced Mannich condensation of terminal alkynes with amines and para-formaldehyde on CuI-doped alumina in the absence of solvents that produces the corresponding aminomethylated adducts in good yields. The reaction can be extended to a Mannich condensation cyclization sequence that generates 2-substituted benzo[b]furans in one-pot and in good yields. The process is highly efficient, does not require pre-forming the iminium species, and is not hampered by the heterogeneity of the reaction medium (Scheme 1).

 $R^1C\equiv CH + (CH_2O)_n + HNR^2R^3 \frac{CuI/AI_2O_3}{MW}$ $R^1C\equiv CCH_2NR^2R^3$

Scheme 1.

Keywords: Cuprous salts; Mannich reaction; Solventles; Microwave.

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Table 1. A Effect of cuprous salts on the Mannich condensation^a

Entry	Cuprous salt	Yield $(\%)^b$
a		28
$\mathbf b$	$Cu2Cl2$ CuBr	26
$\mathbf c$	CuI	82
d	None	

^a Reaction conditions: dibenzylamine (1.00 mmol), 1-decyne (1.00 mmol), *para*-formaldehyde (3.00 mmol), cuprous salt (3.00 mmol), Al_2O_3 (1.00 g), irradiated at 300 W for 10 min.

b Isolated yields.

2. Results and discussion

2.1. Effect of cuprous salts on the Mannich condensation of terminal alkynes with amines and para-formaldehyde

We initially explored the affect of the cuprous salts on the Mannich condensation of terminal alkynes with amines and para-formaldehyde. The results are listed in Table 1. Dibenzylamine, para-formaldehyde and 1-decyne were chosen as the model reactants for this investigation.

It is evident that the Mannich reaction requires a cuprous salt to 'active' the terminal alkyne carbon–hydrogen bond to promote aminomethylation. Among the cuprous salts we

tested, cuprous iodide was most effective and was chosen for further study.

2.2. Mannich condensation of terminal alkynes with secondary amines and *para*-formaldehyde

Table 2 contains a summary of the experimental results. A number of terminal alkynes were successfully condensed with secondary amines and *para*-formaldehyde in good yields. Dibenzylamine, methylbenzylamine, morpholine, piperidine, 1-phenylpiperazine, N-methyl-1-naphthalenemethylamine, di(iso-propyl)amine all reacted smoothly with terminal alkynes and *para*-formaldehyde to generate the corresponding Mannich products. It should be noted that the reaction tolerated many functional groups and that the sterically hindered 2,2,6,6-tetramethylpiperidine smoothly produced the corresponding Mannich adduct.

During the investigation, we found that piperazine reacts with terminal alkynes (2 equiv) and *para*-formaldehyde (excess) to afford the diaminomethylation adducts [bis- (b-aminoalkyne)] ([Scheme 2](#page-88-0)) and 1,9-decadiyne reacted with secondary amines (2 equiv) and *para*-formaldehyde (excess) to generate the bis-Mannich products in moderate to good yields ([Scheme 3](#page-88-0)).

Table 2. Mannich condensation reaction of terminal alkynes with secondary amine and *para-formaldehyde*^a

Entry	Alkyne	Amine	Product	Yield $(\%)^b$
a	$n-C_8H_{17}C\equiv CH$	$(n - C_4H_9)_2NH$	$n-C_8H_{17}C\equiv CCH_2N(n-C_4H_9)_2$	88
$\mathbf b$	n -C ₈ H ₁₇ C \equiv CH	$(C_6H_5CH_2)_2NH$	n -C ₆ H ₁₃ C \equiv CCH ₂ N(CH ₂ CH ₆ H ₅) ₂	82
$\mathbf c$	n -C ₆ H ₁₃ C \equiv CH	$(C6H5CH2)2NH$	n -C ₈ H ₁₇ C \equiv CCH ₂ (CH ₂ C ₆ H ₅) ₂	81
d	n -C ₈ H ₁₇ C=CH	$C_6H_5CH_2NHCH_3$	n -C ₈ H ₁₇ C \equiv CCH ₂ N(CH ₃)CH ₂ C ₆ H ₅	89
$\rm e$	n -C ₈ H ₁₇ C=CH	HN	$n - C_8H_1 \n7C \equiv CCH_2N$	82
f	$n-C_8H_{17}C\equiv CH$	CH ₃ NHCH ₂	n -C ₈ H ₁₇ C \equiv CCH ₂ N(CH ₃)CH ₂	71
g	$C_6H_5C\equiv CH$	$(C_6H_5CH_2)_2NH$	$C_6H_5C\equiv CCH_2N(CH_2C_6H_5)_2$	79
$\mathbf h$	$C_6H_5C\equiv CH$	HN	$C_6H_5C\equiv CCH_2N$	88
\mathbf{i}	p -CH ₃ C ₆ H ₄ C=CH	HN	p -CH ₃ C ₆ H ₄ C \equiv CCH ₂ N	80
j	p -CH ₃ C ₆ H ₄ C=CH	$(C_6H_5CH_2)_2NH$	p -CH ₃ C ₆ H ₄ C \equiv CCH ₂ N(CH ₂ C ₆ H ₅) ₂	83
$\bf k$	p -CH ₃ C ₆ H ₄ C \equiv CH	HN	p -CH ₃ C ₆ H ₄ C \equiv CCH ₂ N	92
$\mathbf{1}$	p -CH ₃ C ₆ H ₄ C \equiv CH	HN	p -CH ₃ C ₆ H ₄ C \equiv CCH ₂ N	74
m	p -FC ₆ H ₄ C=CH	$N - C_6H_5$ HN	p -FC ₆ H ₄ C = CCH ₂ N $N - C_6H_5$	90
$\mathbf n$	p -FC ₆ H ₄ C=CH	$(C_6H_5CH_2)_2NH$	p -FC ₆ H ₄ C \equiv CCH ₂ N(CH ₂ C ₆ H ₅) ₂	85
\mathbf{o}	o -FC ₆ H ₄ C=CH	HN	o -FC ₆ H ₄ C = CCH ₂ N	72
p	p -BrC ₆ H ₄ C=CH	$(n - C_4H_9)$ ₂ NH	$p-\text{BrC}_6\text{H}_4C\equiv \text{CCH}_2\text{N}(n-\text{C}_4\text{H}_9)_{2}$	89

^a Reaction conditions: secondary amine (1.00 mmol), terminal alkyne (1.00 mmol), para-formaldehyde (3.00 mmol), cuprous iodide (3.00 mmol), Al₂O₃ $(1.00 g)$.

b Isolated vields.

Scheme 3.

Scheme 2.

2.3. Mannich condensation of terminal alkynes with primary amines and para-formaldehyde

As anticipated, benzylamine (1 equiv), a primary amine, reacted with 1-decyne (2 equiv) and para-formaldehyde (6 equiv) to produce the bis-Mannich condensation because there are two nitrogen–hydrogen bonds in the primary amine (Scheme 4). Interestingly, when the ratio of reactants is changed alternative reations occur. For example, benzylamine (1 equiv) reacted with 1-decyne (1 equiv) and para-formaldehyde (5 equiv) to generate the Mannich condensation product followed reductive methylation (Scheme 4) phenylacetylene derivatives also afforded methylation products in fair yields (Scheme 4). The reaction provides an alternative route to N -methyl- β -aminoalkynes in a convenient and straightforward fashion.^{[9](#page-96-0)}

2.4. The chemoselectivity of the Mannich condensation of terminal alkynes with secondary amines and para-formaldehyde

The chemoselectivity of the reaction was investigated. When a mixture of acetophenone and 4-ethynyltoluene (or a mixture of 1-decyne and 2-heptanone) served as competitive acidic substrates for the Mannich reaction, only the b-aminoalkynes were formed [\(Scheme 5\)](#page-89-0). As anticipated, the Mannich reaction of 4-acetyl-1-ethynylbenzene with dibenzylamine and para-formaldehyde (or 11-dodecyn-2-one

Scheme 5.

with 1-phenylpiperazine and *para*-formaldehyde) generated b-aminoalkyne products exclusively (Scheme 5).

2.5. 2-Subsitituted benzo[b]furans from the Mannich condensation of o-ethynylphenol with secondary amines and para-formaldehyde

Benzo[b]furans and their derivatives have received much attention in recent years because of their occurrence in natural products and their physiological activity.[10](#page-96-0) They are widely used as antitumor agents, 11 as ligands of the adenosine A_1 receptor,^{[12](#page-96-0)} and as calcium entry blockers.^{[13](#page-96-0)} General routes to benzo $[b]$ furans involve reductive cycliza-tion of ketoesters by low-valent titanium,^{[14](#page-96-0)} photochemically induced rearrangement of phosphate esters, 15 palladium catalyzed Suzuki coupling of boronic acids with organic halides or triflates,^{[16](#page-96-0)} and palladium catalyzed Sonogashira coupling (followed by cyclization) of o -iodophenol and terminal alkynes.[17](#page-96-0) No report has appeared describing the synthesis of 2-substituted benzo[b]furans using a Mannich condensation reaction.

The Mannich condensation–cyclization of o -ethnylphenol with secondary amines and *para*-formaldehyde on cuprous iodide doped alumina under solvent free and microwave irradiation conditions generates 2-(dialkylaminomethyl) benzo[b]furans in good yields (Scheme 6 and [Table 3](#page-90-0)).

[Table 3](#page-90-0) contains a summary of the results. Under microwave irradiation and solvent free conditions, o-ethynylphenol (as well as p -acetyl- o -ethynylphenol) reacts smoothly with para-formaldehyde and a variety of secondary amines, such as 1-phenylpiperizine, piperdine, morpholine, dibutylamine, di(iso-propyl)amine, and N-methylaniline, methylbenzylamine, 1,2,3,4-tetrahydroisoquioine, and N-methylnaphthylmethylamine to afford the desired 2-substituted benzo[b]furans in one-pot. It should be noted that the highly sterically encumbered 2,2,6, 6-tetramethylpiperidine also smoothly undergoes the reaction to generate the corresponding 2-substituted-benzo $[b]$ furan, which was characterized by ${}^{1}H$, ${}^{13}C$ NMR, MS and microanalysis, and confirmed by X-ray crystal diffraction. Interestingly, when o-ethynylphenol (2 equiv) was allowed

Table 3. Mannich condensation–cyclization reaction of o -ethnylphenol and its deriatives with secondary amines and para-formaldehyde (see [Scheme](#page-89-0) [6](#page-89-0)) a

Entry	R	Amine	Yield $\left(\% \right)^{\rm b}$
\rm{a}	H	C_6H_5N NH	65
$\mathbf b$	H	$(n - C_4H_9)_2NH$	68
$\mathbf c$	H	ŃН	65
$\mathrm{d}% \left\ \mathcal{H}\right\ _{A}$	Η	CH ₅ CH ₂ NHCH ₃	62
e	Η	ŅН	55
$\mathbf f$	$\mathbf H$	CH ₂ NHCH ₃	70
g	H	ŃH	59
$\,h$	$\rm H$	$(i$ -C ₃ H ₇) ₂ NH	56
$\rm i$	H	. NH	52
$\mathbf j$	$\rm H$	$C_6H_5NHCH_3$	36
$\bf k$	H	C_6H_5N . NH	65

^a Reaction condition: secondary amine (1.00 mmol), o -ethynylphenol (1.00 mmol), para-formaldehyde (3.00 mmol), cuprous iodide (3.00 mmol), $AI₂O₃$ (1.00 g), irradiated at 300 W for 10 min. b Isolated yields.

to react with para-formaldehyde (excess) and piperazine (1 equiv), a bis-Mannich condensation cyclization product was formed (Scheme 7).

2.6. Surface recyclability

We utilized a surface containing 3 mmol of cuprous iodide per gram of alumina for 1 mmol scale reactions. In an effort to enhance the efficiency of the new solid-state Mannich condensation reaction and reduce waste, recycling was investigated. Table 4 contains a summary of the results. It can be seen that the catalyst and alumina remain active through at least eight cycles. After the product was removed from the surface using an organic solvent, the surface was used directly for the next trial without further treatment.

Table 4. Successive trials for Mannich condensation using $CuI/AI_2O_3^3$

Trial	Yield $(\%)^b$	
	82	
$\overline{2}$	80	
3	81	
$\overline{4}$	79	
5	80	
6	82	
7	80	
8	79	

^a Experiment were carried out as described in the Section 4 by using 1decyne (1 mmol), dibenzylamine (1.00 mmol), para-formaldehyde (3.00 mmol), cuprous iodide (3.00 mmol), Al_2O_3 (1.00 g), microwave irradiation at 300 W for 10 min.

b Isolated yields.

3. Conclusion

A reliable, rapid, practical, and environmentally benign method for the synthesizing β -aminoalkynes and 2-substituted benzo[b]furans has been developed, which involves the use of a solvent-free mixture of cuprous iodide and alumina under microwave irradiation conditions. The process is highly efficient, does not require pre-forming the iminium species, and is not hampered by the heterogeneity of the reaction.

4. Experimental

Melting points were recorded on a MEL-TEMP melting point apparatus and are uncorrected. IR were recorded on a Bomem MB 100 FT-IR. All 1 H and 13 C NMR spectra were recorded on a 250 MHz Bruker AC 250 or Avance 400 MHz spectrometer. Chemical shift are given as δ value with reference to tetramethylsilane (TMS) as internal standard. GC/MS data were obtained by using a Hewlett-Packard 6890 series GC equipped with a 5973 mass selective detector. Microanalyses were performed by Atlantic Microlabs, Norcross, GA. A commercially available Ethos E Touch Control microwave unit (Milestone) was utilized.

Al2O3 and cuprous iodide were purchased from Aldrich Chemical Co. The organic reagents were analytical grade and used as received (Aldrich Chemical Co.).

Products were purified, if applicable, by flash chromatography on 230–400 mesh ASTM 60 Å silica gel, $SiO₂$.

4.1. General procedure for Mannich condensation of terminal alkynes with amines and para-formaldehyde

Secondary amine (1.00 mmol) and terminal alkyne (1.00 mmol) were added to a mixture of cuprous iodide (0.572 g, 3.00 mmol), para-formaldehyde (0.09 g, 3.00 mmol)

and alumina (1.00 g) contained in a clean, dry, 10 mL round-bottomed flask. The mixture was stirred at room temperature to ensure efficient mixing. The flask was then fitted with a septum (punctured by an 18 gauge needle to serve as a pressure release valve), placed in the microwave oven and irradiated at 300 W for 10 min [caution: heating volatile materials in commercial microwave ovens for extended periods can be hazardous]. After cooling, ether (4 mL) was added and the slurry stirred at room temperature to ensure product removal from the surface. The mixture was vacuum filtered using a sintered glass funnel and the product was purified by flash chromatography to yield the desired b-aminoalkyne.

4.1.1. Dibutyl(undec-2-ynyl)amine.^{[18](#page-96-0)} Oil; ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3): \delta \ 3.35 \ (s, 2H), \ 2.38-2.49 \ (t, 4H),$ 2.12–2.21 (t, 3H), 1.19–1.60 (m, 20H), 0.88–0.94 (m, 9H); ¹³C NMR (62.5 MHz, CDCl₃): δ 84.9, 74.7, 53.5, 42.3, 31.7, 29.7, 29.2, 29.1, 29.0, 28.8, 22.7, 20.7, 18.7, 14.1.

4.1.2. Dibenzyl(undec-2-ynyl)amine. Oil; IR (film, $CHCl₃$) 2261 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃) δ 7.40– 7.16 (m, 10H), 3.65 (s, $2 \times 2H$), 3.22 (s, 2H), 2.24 (t, $J=$ 6.6 Hz, 2H), 1.55–1.30 (m, 12H), 0.88 (t, $J=6.8$ Hz, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 139.1, 128.9, 128.1, 126.9, 85.7, 74.4, 57.5, 41.6, 31.8, 29.3, 29.1, 28.9, 22.6, 18.7, 14.1; MS m/z (relative intensity) 347 (M⁺, 3), 270 (6), 256 (10), 194 (7), 91 (100). Anal. Calcd for $C_{25}H_{33}N$: C, 86.40; H, 9.57; N, 4.03. Found: C, 86.34; H, 9.68; N, 4.09.

4.1.3. Dibenzyl(non-2-ynyl)amine. Oil; IR (film, $CHCl₃$) 2260 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.40– 7.18 (m, 10H), 3.65 (s, $2 \times 2H$), 3.22 (s, 2H), 2.25 (t, $J=$ 6.7 Hz, 2H), 1.59–1.31 (m, 8H), 0.91 (t, $J=6.7$ Hz, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 139.1, 129.0, 128.2, 126.9, 85.8, 74.4, 57.5, 41.6, 31.4, 29.1, 28.6, 22.6, 18.7, 14.1; MS m/z (relative intensity) 319 (M⁺, 3), 242 (7), 228 (9), 194 (6), 91 (100). Anal. Calcd for C₂₃H₂₉N: C, 86.47; H, 9.15; N, 4.38. Found: C, 86.24; H, 9.22; N, 4.41.

4.1.4. Benzylmethyl(undec-2-ynyl)amine. Oil; IR (film, CHCl₃) 2261 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.34–7.21 (m, 5H), 3.55 (s, 2H), 3.25 (s, 2H), 2.30 (s, 3H), 2.23 (t, $J=6.8$ Hz, 2H), 1.56–1.28 (m, 12H), 0.88 (t, $J=$ 6.0 Hz, 3H); 13C NMR (62.5 MHz, CDCl3): d 138.6, 129.1, 128.1, 127.0, 85.7, 74.5, 60.1, 45.4, 41.7, 31.8, 29.2, 29.0, 28.9, 28.8, 22.6, 18.7, 14.0; MS m/z (relative intensity) 271 $(M⁺, 7)$, 194 (26), 158 (21), 120 (19), 91 (100). Anal. Calcd for C₁₉H₂₉N: C, 84.07; H, 10.77; N, 5.16. Found: C, 84.21; H, 10.88; N, 5.23.

4.1.5. 1-(Undec-2-ynyl)piperidine. Oil; IR $(\text{film}, \text{CHCl}_3)$ 2188 cm⁻¹ (C \equiv C); ¹H NMR (250 MHz, CDCl₃): δ 3.20 (t, $J=1.8$ Hz, 2H), 2.47 (s, br, $2\times$ 2H), 2.18 (t, $J=6.8$ Hz, 2H), 1.66–1.28 (m, 18H), 0.88 (t, $J=6.5$ Hz, 3H); ¹³C NMR $(62.5 \text{ MHz}, \text{CDCl}_3)$: δ 84.9, 75.0, 53.1, 47.9, 31.6, 29.0, 28.9, 28.7, 25.7, 23.8, 22.4, 18.5, 13.8; MS m/z (relative intensity) 235 (M⁺, 22), 234 (M⁺ -1, 74), 150 (30), 136 (59), 122 (58), 98 (23), 84 (100). Anal. Calcd for $C_{16}H_{29}N$: C, 81.63; H, 12.42; N, 5.95. Found: C, 81.45; H, 12.55; N, 6.02.

4.1.6. Methyl-1-(naphthalenemethyl)(undec-2-ynyl) **amine.** Oil; IR (film, CHCl₃) 2259 cm⁻¹ (C \equiv C); ¹H NMR (250 MHz, CDCl₃): δ 8.29 (d, J = 8.1 Hz, 1H), 7.82– 7.73 (m, 2H), 7.52–7.34 (m, 4H), 3.96 (s, 2H), 3.30 (s, 2H), 2.34 (s, 3H), 2.26 (t, $J=6.7$ Hz, 2H), 1.62–1.28 (m, 12H), 0.87 (t, J=6.6 Hz, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 134.5, 133.8, 132.6, 128.3, 128.0, 127.5, 125.8, 125.5, 125.0, 124.6, 85.9, 74.7, 58.1, 45.6, 41.9, 31.8, 29.2, 29.0, 28.9, 22.6, 18.8, 14.1; MS m/z (relative intensity) 321 (M⁺ 5), 306 (1), 208 (17), 180 (50), 141 (100), 115 (19). Anal. Calcd for $C_{23}H_{31}N$: C, 85.92; H, 9.72; N, 4.36. Found: C, 85.84; H, 9.90; N, 4.44.

4.1.7. Dibenzyl(3-phenylprop-2-ynyl)amine.^{[19](#page-96-0)} Oil; ¹H NMR (250 MHz, CDCl₃): δ 7.45–7.18 (m, 15H), 3.72 (s, 4H), 3.43 (s, 2H); ¹³C NMR (62.5 MHz, CDCl₃): δ 138.8, 131.7, 129.0, 128.3, 127.9, 127.1, 123.4, 85.9, 84.4, 57.9, 42.0.

4.1.8. 1-(3-phenylprop-2-ynyl)piperidine.^{[20](#page-96-0)} Oil; IR (film, CHCl₃) 2188 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.45–7.41 (m, 2H), 7.27–7.25 (m, 3H), 3.46 (s, 2H), 2.55 (s, br, $2 \times 2H$), 1.67–1.59 (m, $2 \times 2H$), 1.45 (t, $J=5.5$ Hz, 2H); ¹³C NMR (62.5 MHz, CDCl₃): δ 131.4, 127.9, 127.7, 123.1, 84.8, 84.7, 53.2, 48.2, 25.7, 23.7; MS m/z (relative intensity) 199 (MC, 42), 170 (11), 157 (25), 130 (7), 122 (13), 115 (100).

4.1.9. 1-[3-(p-Tolyl)prop-2-ynyl]piperidine. 21 21 21 Oil; $^{\mathrm{1}}\mathrm{H}$ NMR (250 MHz, CDCl₃): δ 7.34–7.30 (d, 2H), 7.11–7.07 (d, 2H), 3.46 (s, 2H), 2.50–2.61 (t, 3H), 2.32 (s, 3H), 1.59– 1.70 (t, 4H), 1.36–1.48 (m, 2H); ^{13}C NMR (62.5 MHz, CDCl3): d 137.7, 131.5, 128.8, 120.2, 85.0, 84.2, 53.4, 48.4, 25.9, 23.7, 21.3.

4.1.10. Dibenzyl[3-(p-tolyl)prop-2-ynyl]amine. Oil; IR (film, CHCl₃) 2229 cm^{-1} (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.44–7.20 (m, 12H), 7.11 (d, J=7.9 Hz, 2H), 3.74 (s, $2 \times 2H$), 3.45 (s, 2H), 2.33 (s, 3H); ¹³C NMR (62.5 MHz, CDCl3): d 138.9, 138.0, 131.6, 129.0, 128.2, 127.0, 120.3, 86.0, 83.6, 57.7, 42.0, 21.4; MS m/z (relative intensity) 325 $(M^+, 6)$, 234 (11), 194 (9), 129 (45), 91 (100). Anal. Calcd for C24H23N: C, 88.57; H, 7.12; N, 4.30. Found: C, 88.45; H, 7.21; N, 4.27.

4.1.11. 4-[(3-p-Tolyl)prop-2-ynyl]morpholine. Oil; IR (film, CHCl₃) 2204 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.32 (d, J = 8.1 Hz, 2H), 7.09 (d, J = 8.0 Hz, 2H), 3.75 (t, $J=4.7$ Hz, $2\times 2H$), 3.48 (s, 2H), 2.62 (t, $J=4.6$ Hz, 2×2 H), 2.32 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃); δ 138.0, 131.4, 128.9, 119.8, 85.5, 83.2, 66.7, 52.3, 47.9, 21.3; MS m/z (relative intensity) 215 (M⁺, 29), 184 (32), 170 (26), 157 (40), 129 (100). HRMS Calcd for $C_{14}H_{17}NO$: 215.1310, found: 215.1310.

4.1.12. 2,2,6,6-Tetramethyl-1-[3-(p-tolyl)prop-2-ynyl] piperdine. Mp 178–180 °C; IR (KBr, CHCl₃) 2401 cm⁻ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.27 (d, J=7.9 Hz, 2H), 7.06 (d, J=7.8 Hz, 2H), 3.55 (s, 2H), 2.31 (s, 3H), 1.54–1.44 (m, 6H), 1.17 (s, 12H); ¹³C NMR (62.5 MHz, CDCl3): d 137.3, 131.3, 128.8, 121.2, 92.2, 81.0, 54.9, 41.2, 33.8, 27.5, 21.4, 17.8; MS m/z (relative intensity) 269 (M⁺, 2), 254 (7), 129 (100). Anal. Calcd for $C_{19}H_{27}N$: C, 84.70; H, 10.10; N, 5.20. Found: C, 84.56; H, 10.23; N, 5.11.

4.1.13. 1-[3-(4-Fluorophenyl)prop-2-ynyl]-4-phenylpiperazine. Mp $80.5-81.5$ °C; IR (KBr) 2253 cm⁻ \widehat{C} = C); ¹H NMR (250 MHz, CDCl₃): δ 7.44–7.38 (m, 2H), 7.29–7.23 (m, 2H), 7.01–6.83 (m, 5H), 3.55 (s, 2H), 3.25 (t, $J=4.8$ Hz, $2\times 2H$), 2.78 (t, $J=4.8$ Hz, $2\times 2H$); ¹³C NMR (62.5 MHz, CDCl₃): δ 164.3, 160.3, 151.1, 133.6, 133.4, 129.0, 119.7, 119.0 (d, $J=3.3$ Hz), 116.1, 115.6, 115.2, 84.4, 83.9, 52.0, 49.0, 47.6; MS m/z (relative intensity) 294 (M^+ , 12), 252 (6), 188 (11), 176 (14), 159 (100), 133 (84), 106 (60). Anal. Calcd for C₁₉H₁₉N₂F: C, 77.52; H, 6.51; N, 9.52. Found: C, 77.60; H, 6.66; N, 9.55.

4.1.14. Dibenzyl $[(3-p-fluorophenyl)prop-2-ynyl]$ amine.^{[22](#page-96-0)} Oil; ¹H NMR (250 MHz, CDCl₃): δ 7.44–7.01 (m, 14H), 3.74 (S, 2 \times 2H), 3.46 (s, 2H); ¹³C NMR (62.5 MHz, CDCl3): d 164.3, 160.4, 138.8, 129.9, 129.8, 129.0, 128.3, 127.6, 127.2, 125.2, 118.7, 118.4, 115.5, 115.2, 85.6, 84.7, 57.8, 41.9.

4.1.15. 4-[3-(2-Fluorophenyl)prop-2-ynyl)]morpholine. Oil; IR (film, CHCl₃) 2205 cm^{-1} (C=C); ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3)$: δ 7.46–7.39 (m, 1H), 7.33–7.24 (m, 1H), 7.10–7.02 (m, 2H), 3.77 (t, $J=4.6$ Hz, $2\times$ 2H), 3.56 (s, 2H), 2.66 (s, br, 2 \times 2H); ¹³C NMR (62.5 MHz, CDCl₃): δ 162.8 (d, $J=250.3$ Hz), 133.4, 129.8 (d, $J=7.6$ Hz), 123.8 (d, $J=3.4$ Hz), 115.3 (d, $J=20.8$ Hz), 111.4 (d, $J=$ 15.4 Hz), 89.3, 78.9, 66.8, 52.2, 48.0; MS m/z (relative intensity) 219 (M^+ , 16), 188 (17), 161 (16), 133 (100), 86 (26). Anal. Calcd for $C_{13}H_{14}NFO$: C, 71.21; H, 6.44; N, 6.39. Found: C, 70.93; H, 6.50; N, 6.28.

4.1.16. [3-(4-Bromophenyl)prop-2-ynyl]dibutylamine. Oil; IR (film, CHCl₃) 2190 cm^{-1} (C=C); ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3)$: δ 7.41 (d, J=8.5 Hz, 2H), 7.27 (d, $J=8.5$ Hz, 2H), 3.58 (s, 2H), 2.51 (t, $J=7.3$ Hz, $2\times$ 2H), 1.51–1.27 (m, 8H), 0.93 (t, $J=7.1$ Hz, 2×3 H); ¹³C NMR (62.5 MHz, CDCl3): d 133.0, 131.3, 122.3, 121.9, 86.2, 83.8, 53.6, 42.6, 29.6, 20.6, 14.0; MS m/z (relative intensity) $323, 321 \ (M^+, 4, 4), 280, 278 \ (97, 100), 195, 193 \ (78, 79),$ 114 (26), 84 (20). Anal. Calcd for $C_{17}H_{24}NBr: C$, 63.36; H, 7.51; N, 4.35. Found: C, 63.31; H, 7.53; N, 4.37.

4.2. General procedure for the Mannich condensation of terminal alkyne with piperazine and para-formaldehyde

Terminal alkyne (2.00 mmol), para-formaldehyde (6.00 mmol) and piperazine (1.00 mmol) were mixed well with $A1_2O_3$ (2.00 g) and cuprous iodide (1.16 g, 6.00 mmol) and placed in the microwave oven and irradiated at 30% power for 10 min.

4.2.1. 1,4-Di(non-2-ynyl)piperazine. Oil; IR (film, $CHCl₃$) 2196 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): 3.25 (s, $2 \times 2H$), 2.62 (s, br, $4 \times 2H$), 2.17 (t, $J=6.9$ Hz, $2 \times 2H$), 1.52–1.26 (m, 16H), 0.89 (t, $J=6.7$ Hz, 2 \times 3H); ¹³C NMR $(62.5 \text{ MHz}, \text{CDCl}_3)$: δ 85.3, 74.5, 51.7, 47.1, 31.1, 28.6, 28.4, 22.3, 18.5, 13.8; MS m/z (relative intensity) 330 (M⁺, 3), 287 (10), 260 (9), 231 (10), 207 (100), 178 (26), 108 (20). Anal. Calcd for $C_{22}H_{38}N_2$: C, 79.94; H, 11.59; N, 8.47. Found: C, 76.69; H, 11.68; N, 8.43.

4.2.2. 1,4-Di[3-(p-tolyl)prop-2-ynyl]piperazine. Mp 119– 121 °C; IR (film, CHCl₃) 2199 cm⁻¹ (C=C); ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3)$ 7.32 (d, $J=8.0 \text{ Hz}, 2 \times 2\text{H}$), 7.08 (d, $J=$ 8.1 Hz, $2 \times 2H$), 3.52 (s, $2 \times 2H$), 2.74 (s, br, $2 \times 4H$), 2.32 (s, 2 \times 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 138.0, 131.5, 128.9, 120.0, 85.4, 83.5, 51.9, 47.6, 21.3; MS m/z (relative intensity) 342 (M⁺, 2), 341 (M⁺ -1, 5), 272 (3), 223 (5), 213 (11), 129 (100), 115 (8). Anal. Calcd for $C_{24}H_{26}N_2$: C, 84.17; H, 7.65; N, 8.18. Found: C, 83.96; H, 7.85; N, 8.03.

4.3. General procedure for the Mannich condensation of α, ω -dialkyne with secondary amine and para-formaldehyde

 α , ω -Dialkyne (1.00 mmol), para-formaldehyde (6.00 mmol) and secondary amine (2.00 mmol) were mixed well with Al₂O₃ (2.00 g) and cuprous iodide (1.16 g, 6.00 mmol) and placed in the microwave oven and irradiated at 300 W for 10 min.

4.3.1. N,N,N,N-Tetrabutyl-2,10-dodecadiynyl-1,12 **diamine.** Oil; IR (film, CHCl₃) 2189 cm^{-1} (C \equiv C); ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3)$ 3.34 (t, $J=1.9 \text{ Hz}, 2 \times 2\text{H}$), 2.43 (t, $J=$ 7.3 Hz, $4 \times 2H$), 2.19 (t, $J=6.5$ Hz, $2 \times 2H$), 1.53–1.25 (m, 24H), 0.92 (t, $J=7.1$ Hz, 4×3 H); ¹³C NMR (62.5 MHz, CDCl3): d 84.5, 74.7, 53.3, 42.0, 29.5, 28.7, 28.1, 20.6, 18.5, 13.9; MS m/z (relative intensity) 246 (M⁺ -170, 21), 112 (12), 84 (100), 70 (29), 57 (66). Anal. Calcd for $C_{28}H_{52}N_2$: C, 80.70; H, 12.58; N, 6.72. Found: C, 80.50; H, 12.62; N, 6.69.

4.3.2. 1,12-Di(4-phenylpiperazino)-2,10-dodecadiyne. Mp 81–82 °C; IR (film, CHCl₃) 2204 cm⁻¹ (C \equiv C); ¹H NMR (250 MHz, CDCl₃) 7.28–7.21 (m, $2 \times 2H$), 6.93–6.81 $(m, 2 \times 3H), 3.29$ (s, 2 \times 2H), 3.22 (t, J=7.7 Hz, 4 \times 2H), 2.69 (t, $J=4.6$ Hz, $4 \times 2H$), 2.20 (t, $J=6.8$ Hz, $2 \times 2H$), 1.53–1.39 (m, 8H); ¹³C NMR (62.5 MHz, CDCl₃): δ 151.1, 128.9, 119.5, 85.4, 74.5, 51.9, 48.9, 47.2, 28.5, 28.1, 18.5; MS m/z (relative intensity) 322 (M⁺ -160, 17), 216 (10), 159 (94), 120 (66), 106 (100), 77 (70). Anal. Calcd for $C_{32}H_{42}N_{4}$: C, 79.62; H, 8.77; N, 11.61. Found: C, 79.60; H, 8.76; N, 11.55.

4.4. Mannich condensation reaction of terminal alkyne with primayl amine and para-formaldehyde

(a) 1-Decyne (2.00 mmol), para-formaldehyde (6.00 mmol) and benzylamine (2.00 mmol) were mixed well with Al_2O_3 (2.00 g) and cuprous iodide (1.16 g, 6.00 mmol) and placed in the microwave oven and irradiated at 300 W for 10 min, worked up the same as the general procedure. (b) Terminal alkyne (1.00 mmol), para-formaldehyde (5.00 mmol) and benzylamine (1.00 mmol) were mixed well with Al_2O_3 (1.00 g) and cuprous iodide (0.58 g, 3.00 mmol) and placed in the microwave oven and irradiated at 30% power for 10 min.

4.4.1. Benzyldi(undec-2-ynyl)amine. Oil; IR (film, $CHCl₃$) 2232 cm⁻¹ (C \equiv C); ¹H NMR (250 MHz, CDCl₃) 7.38–7.20 (m, 5H), 3.66 (s, 2H), 3.35 (s, $2 \times 2H$), 2.21 (t, $J=6.7$ Hz, 2×2 H), 1.55–1.28 (m, 2 \times 12H), 0.88 (t, J=6.6 Hz, 2 \times 3H); 13C NMR (62.5 MHz, CDCl3): d 138.3, 129.3, 128.1, 127.1, 85.3, 75.0, 56.9, 42.3, 31.8, 29.2, 29.1, 28.9, 22.6,

18.7, 14.0; MS m/z (relative intensity) 322 (M⁺ -85, 6), 294 (7), 184 (5), 156 (8), 91 (100). Anal. Calcd for C29H45N: C, 85.44; H, 11.13; N, 3.44. Found: C, 85.37; H, 11.24; N, 3.52.

4.4.2. Benzylmethyl(undec-2-ynyl)amine. Oil; IR (film, CHCl₃) 2260 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl₃): δ 7.35–7.22 (m, 5H), 3.55 (s, 2H), 3.26 (t, $J=2.0$ Hz, 2H), 2.30 (s, 3H), 2.26–2.20 (m, 2H), 1.59–1.28 (m, 12H), 0.88 (t, $J=6.5$ Hz, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 138.6, 129.2, 128.2, 127.1, 85.8, 74.6, 60.2, 45.5, 41.8, 31.8, 29.2, 29.1, 29.0, 28.9, 22.6, 18.7, 14.0; MS m/z (relative intensity) 271 (M⁺, 7), 194 (25), 158 (22), 120 (18), 91 (100). Anal. Calcd for $C_{19}H_{29}N$: C, 84.07; H, 10.77; N, 5.16. Found: C, 83.96; H, 10.85; N, 5.19.

4.4.3. Benzylmethyl[3-(phenyl)prop-2-ynyl]amine. Oil; IR (film, CHCl₃) 2235 cm⁻¹ (C=C); ¹H NMR (250 MHz, CDCl3): d 7.48–7.44 (m, 2H), 7.38–7.23 (m, 8H), 3.62 (s, 2H), 3.49 (s, 2H), 2.38 (s, 3H); 13C NMR (62.5 MHz, CDCl3): d 138.3, 131.6, 129.1, 128.1, 127.9, 127.1, 123.2, 85.6, 84.3, 60.1, 45.6, 41.8; MS m/z (relative intensity) 235 (M^+ , 18), 158 (41), 144 (27), 115 (100), 91 (64). Anal. Calcd for $C_{17}H_{17}N$: C, 86.77; H, 7.28; N, 5.95. Found: C, 86.66; H, 7.43; N, 6.01.

4.4.4. Benzylmethyl[3-(p-tolyl)prop-2-ynyl]amine. Oil; IR (film, CHCl₃) 2247 cm^{-1} (C=C); ¹H NMR $(250 \text{ MHz}, \text{ CDC1}_3)$: δ 7.37–7.24 (m, 7H), 7.09 (d, J= 7.94 Hz, 2H), 3.62 (s, 2H), 3.49 (s, 2H), 2.38 (s, 3H), 2.32 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 138.4, 137.9, 131.5, 129.1, 128.9, 128.2, 127.1, 120.2, 85.7, 83.6, 60.2, 45.7, 41.9, 21.3; MS m/z (relative intensity) 249 (M⁺, 17), 172 (32) , 158 (36), 129 (100), 91 (73). Anal. Calcd for $C_{18}H_{19}N$: C, 86.70; H, 7.68; N, 5.62. Found: C, 86.42; H, 7.60; N, 5.57.

4.4.5. Preparation of 4-acetyl-1-ethynylbenzene. 23 23 23 To a stirred solution of p -bromoacetophone (1.00 g, 5.00 mmol) and Et₃N–dioxane (4 mL/4 mL), PdCl₂(PPh₃)₂ (0.035 g, 0.05 mmol) and CuI (0.019 g, 0.100 mmol) were added in one portion. Then (trimethylsilyl)acetylene (0.85 mL, 6.00 mmol) was added dropwise at room temperature under nitrogen. The reaction mixture was stirred overnight. Et3N, dioxane and unreacted (trimethylsilyl)acetylene were removed under reduced pressure. The product extracted with $Et₂O$ (3 × 20 mL). The combined organic phase was washed with H_2O , dried (Na₂SO₄) and concentrated under vacuum. The residue was purified by column chromatography to generate 0.820 g (85% yield) of 4-acetyl-1- ((trimethylsilyl)ethynyl)benzene. Oil; ¹ H NMR (250 MHz, CDCl₃): δ 7.87 (d, J=7.9 Hz, 2H), 7.52 (d, J=8.4 Hz, 2H), 2.57 (s, 3H), 0.27 (s, 9H); ¹³C NMR (62.5 MHz, CDCl₃): δ 196.9, 136.3, 131.9, 128.0, 127.8, 103.9, 97.9, 26.4, -0.29; MS m/z (relative intensity) 216 (M⁺, 15), 201 (100), 158 (10), 143 (8), 93 (9).

4-Acetyl-1-((trimethylsilyl)ethynyl)benzene (0.648 g, 3.00 mmol) was added to KF/Al_2O_3 (2.00 g, 40% by weight) and stirred at room temperature to ensure efficient mixing. The result mixture was placed in the microwave oven and irradiated at 30% power for 3 min. After cooling, hexane (10 mL) was added and the slurry stirred at room

temperature to ensure product removal from the surface. The product was purified by chromatography to afford 0.415 g (97% yield) of 4-acetyl-1-ethynylbenzene. Mp 68– 70 °C (Iit.^{[23](#page-96-0)} 69–70 °C); ¹H NMR (250 MHz, CDCl₃): δ 7.90 (d, $J=8.3$ Hz, 2H), 7.57 (d, $J=8.4$ Hz, 2H), 3.27 (s, 1H), 2.59 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 197.1, 136.7, 132.2, 128.1, 126.8, 82.7, 80.3, 26.5; MS m/z (relative intensity) 144 (M^+ , 31), 129 (100), 101 (54), 75 (21).

4.4.6. Preparation of 11-dodecyn-2-one.^{[24](#page-96-0)} To a suspension of dry pyridinium chlorochromate (4.85 g, 22.5 mmol) in dry CH_2Cl_2 (20 mL), 10-undecyn-1-ol (2.50 g, 15 mmol) and dry CH_2Cl_2 (5 mL) were added dropwise. The reaction mixture was stirred at room temperature for 3 h and the solution was then extracted with ether $(3 \times 20 \text{ mL})$. The combined organic layer was dried with $Na₂SO₄$ and the solvent removed under reduced pressure. The product was purified by column chromatography (hexane/ethyl acetate 9:1) to afford 2.00 g (80% yield) of 10-undecyn-1-al.

To a solution of 10-undecyn-1-al (1.50 g, 9.04 mmol) in dry Et₂O (15 mL), CH₃MgBr in ether (3 M, 3.67 mL, 11 mmol) was added at 0° C over a 1 h period. The reaction mixture was then refluxed for 3 h. Aqueous ammonium chloride was then added to quench the reaction and the mixture was extracted with ether $(3 \times 20 \text{ mL})$. The combined organic layer was washed with water and dried with $Na₂SO₄$ and then the solvent removed under reduced pressure (2 mmHg for 2 h). The crude product was used without purification.

11-Dodecyn-2-ol (crude, 1.4 g) in dry $CH₂Cl₂$ (5 mL) was added dropwise to a suspension of pyridinium chlorochromate (2.26 g, 10.5 mmol) in dry CH_2Cl_2 (10 mL) dropwise at room temperature. The reaction mixture was stirred at room temperature for 2 days and then extracted with ether $(3 \times 20 \text{ mL})$. The combined organic layer was dried with $Na₂SO₄$ and then the solvent removed under reduced pressure. The product was purified by column chromatography (hexane/ethyl acetate 85:15) to afford 1.20 g (combined yield 74%) of 11-dodecyn-2-one. Oil; ¹H NMR $(250 \text{ MHz}, \text{CDCl}_3)$: δ 2.41 (t, J = 7.4 Hz, 2H), 2.18–2.12 (m, 5H), 1.93 (t, $J=2.4$ Hz, 1H), 1.55–1.28 (m, 12H); ¹³C NMR $(62.5 \text{ MHz}, \text{CDCl}_3)$: δ 208.5, 84.2, 67.9, 43.4, 29.5, 28.9, 28.8, 28.6, 28.3, 28.1, 23.5, 18.0; MS m/z (relative intensity) $165 \ (M^+-C_2H_5, 2), 147 \ (2), 122 \ (6), 107 \ (9), 95 \ (15), 81$ (23), 58 (100).

4.5. Chemoselectivity of Mannich Condensation

(a) Intermolecular chemoselectivity. A secondary amine (morpholine or piperazine, 1.00 mmol), terminal alkyne (4-ethynyltoluene or 1-decyne, 1.00 mmol) and an enolizable ketone (acetophone or 2-heptone, 1.00 mmol) were mixed with a mixture of cuprous iodide (0.572 g, 3.00 mmol), para-formaldehyde (0.09 g, 3.00 mmol) and alumina (1.00 g) in a clean, dry, 10 mL round-bottomed flask at room temperature. The flask was then fitted with a septum (punctured by an 18 gauge needle), placed in the microwave oven and irradiated at 30% power for 10 min.

(b) Intramolecular chemoselectivity. Dibenzylamine or 1-phenylpiperazine (1.00 mmol) and 4-acetyl-1-ethynylbenzene or 11-dodecyn-2-one (1.00 mmol) were added to a mixture of cuprous iodide (0.572 g, 3.00 mmol), paraformaldehyde $(0.09 \text{ g}, 3.00 \text{ mmol})$ and alumina (1.00 g) contained in a clean, dry, 10 mL round-bottomed flask. The mixture was stirred at room temperature to ensure efficient mixing. The flask was then fitted with a septum (punctured by an 18 gauge needle), placed in the microwave oven and irradiated at 30% power for 10 min.

4.5.1. [3-(4-Acetylphenyl)prop-2-ynyl]dibenzylamine. Oil; IR (film, CHCl₃) 2230 cm⁻¹ (C=C), 1683 cm⁻ (C=O); ¹H NMR (250 MHz, CDCl₃): δ 7.92 (d, J = 8.3 Hz, 2H), 7.56 (d, $J=8.2$ Hz, 2H), 7.45–7.23 (m, 10H), 3.76 (s, $2 \times 2H$), 3.50 (s, 2H), 2.60 (s, 3H); ¹³C NMR (62.5 MHz, CDCl3): d 197.2, 138.7, 136.1, 131.9, 129.0, 128.3, 128.2, 127.2, 88.3, 85.2, 57.8, 42.1, 26.6; MS m/z (relative intensity) 194 ($M⁺ - 159$, 11), 158 (15), 143 (25), 115 (20), 91 (100). Anal. Calcd for $C_{25}H_{23}NO: C$, 84.95; H, 6.56; N, 3.96. Found: C, 85.11; H, 6.66; N, 3.94.

4.5.2. 1-Phenyl-4-(tridec-12-oxo-2-ynyl)piperazine. Mp 33–34 °C; IR (film, CHCl₃) 2208 cm⁻¹ (C=C), 1711 cm⁻¹ (C=O); ¹H NMR (250 MHz, CDCl₃): δ 7.29-7.22 (m, 2H), $6.95-6.82$ (m, 3H), 3.31 (s, 2H), 3.23 (t, $J=$ 4.9 Hz, $2 \times 2H$), 2.71 (t, $J=4.9$ Hz, $2 \times 2H$), 2.39 (t, $J=$ 7.4 Hz, 2H), 2.19 (t, $J=6.7$ Hz, 2H), 2.11 (s, 3H), 1.58–1.28 (m, 12H); ¹³C NMR (100 MHz, CDCl₃): δ 209.2, 151.1, 129.0, 119.6, 116.0, 85.6, 74.5, 51.9, 48.9, 47.3, 43.6, 29.7, 29.2, 29.0, 28.8, 28.7, 28.6, 23.7, 18.6; MS m/z (relative intensity) 354 (M^+ , 18), 248 (35), 160 (53), 159 (94), 130 (38), 120 (71), 105 (100). Anal. Calcd for $C_{23}H_{34}N_2O$: C, 77.92; H, 9.67; N, 7.90. Found: C, 77.98; H, 9.61; N, 8.02.

4.5.3. Iodination of phenols: preparation of 4-hydroxy-3 iodoacetophenone (representative procedure). 4-Hydroxyacetophenone (1.36 g, 10 mmol) was dissolved in 10 mL of THF–H₂O (50/50, V/V) and I_2 (2.80 g, 11 mmol) and NaHCO₃ (0.92 g, 11 mmol) were crushed together and added to the solution. After the mixture was stirred for 3 h at room temperature, residual I_2 was destroyed by addition of a 5% aqueous solution of $Na₂S₂O₃$ until the brown color disappeared. The mixture was extracted with ether $(3) \times$ 50 mL). The organic phase was dried with $Na₂SO₄$ and the solvent was removed under pressure. 4-Hydroxy-3-iodoacetophenone (1.23 g, 47% yield) was isolated by column chromatography (silica gel, hexane/ethyl acetate 95:5). Mp $152-154$ °C (lit.^{[25](#page-96-0)} 153-155 °C); ¹H NMR (250 MHz, CD₃OD): δ 8.29 (s, 1H), 7.82 (d, J=8.5 Hz, 1H), 6.86 (d, $J=8.4$ Hz, 1H), 4.91 (s, br, 1H), 2.51 (s, 3H); MS m/z (relative intensity) 262 (M^+ , 36), 247 (100), 219 (16), 127 (7), 120 (23), 92 (40).

4.5.4. Preparation of o-ethynylphenol and p-acetyl-oethynylphenol via a Sonogashira coupling followed by desilylation: synthesis of o-ethynylphenol as representative. To a stirred solution of 2-iodophenol (1.10 g, 5.00 mmol) and Et₃N–dioxane (4 mL/4 mL), PdCl₂(PPh₃)₂ (0.035 g, 0.05 mmol) and CuI (0.019 g, 0.10 mmol) were added in one portion. Then (trimethylsilyl)acetylene (0.85 mL, 6.00 mmol) was added dropwise at room temperature under nitrogen. The reaction mixture was stirred overnight. Et₃N, dioxane and unreacted (trimethylsilyl)acetylene were removed under reduced pressure. The product was extracted with Et_2O (3×20 mL) and the

combined organic phase washed with H_2O , dried (Na_2SO_4) and concentrated under vacuum. The residue was purified by column chromatography to afford 0.87 g (86% yield) of o -((trimethylsilyl)ethynyl)phenol. Mp 46–48 °C (lit.^{[26](#page-96-0)}) 46–47 °C); ¹H NMR (250 MHz, CDCl₃): δ 7.32 (dd, J= 6.3 Hz, $J=1.4$ Hz, 1H), 7.20 (dt, $J=7.0$ Hz, $J=1.5$ Hz, 1H), 6.93 (d, $J=8.2$ Hz, 1H), 6.82 (t, $J=7.6$ Hz, 1H), 5.90 (s, 1H), 0.26 (s, 9H); ¹³C NMR (62.5 MHz, CDCl₃): d 157.0, 131.6, 130.6, 120.1, 114.5, 109.5, 102.2, 99.0, -0.1 ; MS *m/z* (relative intensity) 190 (M⁺, 18), 175 (100), 159 (19), 135 (13), 115 (16), 77 (14).

o-[(Trimethylsilyl)ethynyl]phenol (0.61 g, 3.00 mmol) was added to KF/Al_2O_3 (2.00 g, 40% by weight) and stirred at room temperature to ensure efficient mixing. The resultant mixture was placed in the microwave oven and irradiated at 30% power for 3 min. After cooling, hexane (10 mL) was added and the slurry stirred at room temperature to ensure product removal from the surface. The product was purified by chromatography to afford 0.365 g (92% yield) of o -ethynylphenol. Oil;^{[27](#page-96-0)}¹H NMR (250 MHz, CDCl₃): δ 7.38 (d, $J=7.5$ Hz, 1H), 7.27 (dt, $J=7.7$ Hz, $J=1.0$ Hz, 1H), 6.95 (d, $J=8.2$ Hz, 1H), 6.87 (t, $J=7.5$ Hz, 1H), 5.80 (s, br, 1H), 3.46 (s, 1H); ¹³C NMR (62.5 MHz, CDCl₃): δ 157.4, 132.0, 130.9, 120.3, 114.8, 108.3, 84.3, 78.3; MS m/z (relative intensity) 118 (M^+ , 100), 89 (44), 63 (21).

4.5.5. p-Acetyl-o-[(trimethylsilyl)ethynyl]phenol. Mp 127–129 °C; ¹H NMR (250 MHz, CDCl₃): δ 8.00 (d, J= 1.4 Hz, 1H), 7.87 (d, $J=8.7$ Hz, 1H), 7.00 (d, $J=8.7$ Hz, 1H), 6.63 (s, br, 1H), 2.55 (s, 3H), 0.29 (s, 9H); 13C NMR (62.5 MHz, CDCl3): d 196.2, 160.9, 133.0, 131.1, 129.9, 114.8, 109.9, 103.2, 97.8, 26.2, -0.2 ; MS m/z (relative intensity) 232 (M^+ , 24), 217 (100), 174 (8), 115 (10), 101 (19). Anal. Calcd for $C_{13}H_{16}O_2Si$: C, 67.20; H, 6.94. Found: C, 67.40; H, 6.84.

4.5.6. p-Acetyl-o-ethynylphenol. Mp $100-102$ °C; 1 H NMR (250 MHz, CDCl₃): δ 8.05 (d, J = 1.5 Hz, 1H), 7.91 (d, $J=8.7$ Hz, 1H), 7.37 (s, 1H), 7.03 (d, $J=8.6$ Hz, 1H), 3.50 (s, 1H), 2.57 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 196.7, 161.6, 133.8, 131.3, 129.6, 115.2, 108.8, 84.4, 77.5, 26.1; MS m/z (relative intensity) 160 (M⁺, 40), 145 (100), 117 (48), 89 (31). HRMS Calcd for $C_{10}H_8O_2$ 160.0524, found 160.0520.

4.6. One-pot synthesis of 2-substitutited benzo $[b]$ furans via the Mannich condensation cyclization sequence reaction of o-ethynylphenol with secondary amines and para-formaldehyde: general procedure

o-Ethynylphenol (0.18 g, 1.00 mmol) and the secondary amine (1.00 mmol) were added to a mixture of cuprous iodide (0.57 g, 3.00 mmol), para-formaldehyde (0.09 g, 3.00 mmol) and alumina (1.00 g) contained in a clean, dry, 10 mL round-bottomed flask. The mixture was stirred at room temperature to ensure efficient mixing. The flask was then fitted with a septum (punctured by an 18 gauge needle), placed in the microwave oven and irradiated at 30% power for 10 min. After cooling, ether (4 mL) was added and the slurry stirred at room temperature to ensure product removal from the surface. The mixture was vacuum filtered using a sintered glass funnel and the product purified by flash chromatography (hexane/EtOAc as eluting agent) to afford the desired 2-substituted-benzo $[b]$ furan.

4.6.1. 1-(Benzofuran-2-ylmethyl)-4-phenylpiperazine. Mp 75–76 °C; ¹H NMR (250 MHz, CDCl₃): δ 7.53–7.46 (m, 2H), 7.26–7.18 (m, 4H), 6.90–6.80 (m, 3H), 6.59 (s, 1H), 3.70 (s, 2H), 3.20 (t, $J=4.8$ Hz, $2\times$ 2H), 2.67 (t, $J=$ 4.8 Hz, 2×2 H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.0, 154.2, 151.1, 129.0, 128.1, 123.9, 122.6, 120.6, 119.6, 116.0, 111.2, 105.7, 55.3, 52.9, 48.9; MS m/z (relative intensity) 292 (M⁺, 33), 186 (12), 173 (22), 159 (98), 131 (100), 106 (54), 77 (56). Anal. Calcd for $C_{19}H_{20}N_2O$: C, 78.05; H, 6.89; N, 9.58. Found: C, 78.04; H, 6.89; N, 9.47.

4.6.2. (Benzofuran-2-ylmethyl)dibutylamine.^{[28](#page-96-0)} Oil; ¹H NMR (250 MHz, CDCl₃): δ 7.50–7.43 (m, 2H), 7.22–7.17 (m, 2H), 6.54 (s, 1H), 3.77 (s, 2H), 2.50 (t, $J=7.5$ Hz, 2 \times 2H), 1.53–1.46 (m, 2×2 H), 1.35–1.27 (m, 2×2 H), 0.90 (t, $J=7.2$ Hz, $2\times3H$); ¹³C NMR (62.5 MHz, CDCl₃): d 156.2, 154.9, 128.5, 123.5, 122.4, 120.5, 111.1, 104.8, 53.7, 50.7, 29.2, 20.6, 14.0; MS m/z (relative intensity) 259 $(M⁺, 2), 216 (10), 131 (100), 77 (9).$

4.6.3. 1-(Benzofuran-2-ylmethyl)piperidine.^{[29](#page-96-0)} Oil; ¹H NMR (250 MHz, CDCl₃): δ 7.53–7.45 (m, 2H), 7.27–7.15 $(m, 2H)$, 6.56 (s, 1H), 3.64 (s, 2H), 2.46 (s, br, 2×2H), 1.63–1.57 (m, $2 \times 2H$), 1.44–1.42 (m, 2H); ¹³C NMR (62.5 MHz, CDCl3): d 155.0, 128.3, 123.7, 122.5, 120.5, 111.2, 105.3, 56.1, 54.3, 25.8, 24.1; MS m/z (relative intensity) 251 (M^+ , 10), 131 (100), 84 (26), 77 (15).

4.6.4. (Benzofuran-2-ylmethyl)benzylmethylamine. Oil; ¹ ¹H NMR (250 MHz, CDCl₃): δ 7.52–7.18 (m, 9H), 6.57 (s, 1H), 3.70 (s, 2H), 3.60 (s, 2H), 2.29 (s, 3H); 13C NMR (62.5 MHz, CDCl3): d 155.4, 155.0, 138.4, 129.0, 128.3, 128.2, 127.1, 123.7, 122.5, 120.6, 111.2, 105.2, 61.3, 53.7, 42.1; MS m/z (relative intensity) 251 (M⁺, 10), 160 (13), 131 (100), 91 (27), 77 (13). Anal. Calcd for $C_{17}H_{17}NO: C$, 81.24; H, 6.82; N, 5.57. Found: C, 81.25; H, 6.82; N, 5.56.

4.6.5. 4-(Benzofuran-2-ylmethyl)morpholine. Mp 50– 51 °C (lit.^{[30](#page-96-0)} 51–52 °C); ¹H NMR (250 MHz, CDCl₃): δ 7.54–7.46 (m, 2H), 7.28–7.17 (m, 2H), 6.60 (s, 1H), 3.74 (t, $J=4.6$ Hz, $2\times$ 2H), 3.67 (s, 2H), 2.54 (t, $J=4.5$ Hz, 2 \times 2H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.1, 154.0, 128.1, 123.9, 122.6, 120.6, 111.2, 105.8, 66.7, 55.8, 53.4; MS m/z (relative intensity) 217 (M^+ , 14), 144 (9), 131 (100), 86 (26), 77 (17).

4.6.6. (Benzofuran-2-ylmethyl)methyl(1-naphthalenemethyl)amine. Oil; ¹H NMR (250 MHz, CDCl₃): δ 8.20– 8.17 (m, 1H), 7.80–7.70 (m, 2H), 7.50–7.32 (m, 6H), 7.26– 7.14 (m, 2H), 6.57 (s, 1H), 3.94 (s, 2H), 3.76 (s, 2H), 2.30 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.5, 155.0, 134.3, 133.8, 132.4, 128.3, 128.0, 127.4, 125.9, 125.5, 125.0, 124.5, 123.8, 122.5, 120.6, 111.1, 105.4, 59.4, 54.2, 42.3; MS m/z (relative intensity) 301 (M^+ , 9), 207 (8), 160 (42), 141 (61), 131 (100), 115 (19), 77 (13). Anal. Calcd for $C_{21}H_{19}NO: C$, 83.69; H, 6.35; N, 4.65. Found: C, 83.71; H, 6.53; N, 4.68.

4.6.7. 2-(Benzofuran-2-ylmethyl)-1,2,3,4-tetrahydro**isoquinoline.** Mp 58–59 °C; ¹H NMR (250 MHz, CDCl₃): δ 7.53–7.45 (m, 2H), 7.25–6.95 (m, 6H), 6.62 (s, 1H), 3.83 $(s, 2H), 3.71 (s, 2H), 2.90 (t, J=5.3 Hz, 2H), 2.82 (t, J=$ 5.4 Hz, 2H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.0, 154.6, 134.3, 133.9, 128.5, 128.2, 126.5, 126.1, 125.5, 123.8, 122.5, 120.6, 111.2, 105.4, 55.6, 54.9, 50.5, 28.8; MS m/z (relative intensity) 263 (M^+ , 5), 145 (19), 131 (100), 104 (23), 77 (21). Anal. Calcd for $C_{18}H_{17}NO$: C, 82.10; H, 6.51; N, 5.32. Found: C, 81.96; H, 6.57; N, 5.30.

4.6.8. (Benzofuran-2-ylmethyl)di(isopropyl)amine. Oil; ¹ ¹H NMR (250 MHz, CDCl₃): δ 7.50–7.41 (m, 2H), 7.20– 7.16 (m, 2H), 6.57 (s, 1H), 3.77 (d, $J=0.59$ Hz, 2H), 3.18– 3.08 (m, 2H), 1.06 (d, $J=6.6$ Hz, 4×3 H); ^{13}C NMR (62.5 MHz, CDCl3): d 160.2, 154.8, 128.9, 123.0, 122.3, 120.3, 110.9, 103.2, 49.0, 42.9, 20.7; MS m/z (relative intensity) 231 (M^+ , 3), 216 (9), 131 (100). Anal. Calcd for $C_{15}H_{21}NO: C, 77.88; H, 9.15; N, 6.05. Found: C, 78.16; H,$ 9.36; N, 5.97.

4.6.9. 1-(Benzofuran-2-ylmethyl)-2,2,6,6-tetramethylpiperidine. Mp 88–89 °C; ¹H NMR (250 MHz, CDCl₃): δ 7.48–7.36 (m, 2H), 7.17–7.14 (m, 2H), 6.63 (s, 1H), 3.81 (s, 2H), 1.60–1.51 (m, 6H), 1.06 (s, 4×3 H); ¹³C NMR (62.5 MHz, CDCl3): d 163.5, 154.5, 129.1, 122.7, 122.2, 120.2, 110.7, 102.9, 54.9, 42.4, 41.2, 27.4, 17.8; MS m/z (relative intensity) 271 (M^+ , 2), 256 (15), 131 (100), 77 (7). Anal. Calcd for $C_{18}H_{25}NO$: C, 79.66; H, 9.28; N, 5.16. Found: C, 79.40; H, 9.36; N, 4.91.

4.6.10. (Benzofuran-2-ylmethyl)methylaniline. Oil; ${}^{1}H$ NMR (250 MHz, CDCl₃): δ 7.48–7.41 (m, 2H), 7.28–7.15 (m, 4H), 6.86–6.72 (m, 3H), 6.49 (s, 1H), 4.61 (s, 2H), 3.08 (s, 3H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.2, 154.9, 149.1, 129.2, 128.4, 123.7, 122.6, 120.6, 117.3, 112.9, 111.0, 103.9, 50.5, 38.6; MS m/z (relative intensity) 237 $(M⁺, 13)$, 207 (26), 131 (100), 96 (9), 77 (23). Anal. Calcd for $C_{16}H_{15}NO$: C, 80.98; H, 6.37; N, 5.90. Found: C, 81.17; H, 6.61; N, 5.63.

4.6.11. 1-(5-Acetylbenzofuran-2-ylmethyl)-4-phenyl**piperazine.** Mp 111–113 °C; ¹H NMR (250 MHz, CDCl₃): δ 8.19 (s, 1H), 7.93 (d, J = 8.6 Hz, 1H), 7.52 (d, J = 8.7 Hz, 1H), 7.25 (t, $J=7.7$ Hz, 2H), 6.93–6.82 (m, 3H), 6.17 (s, 1H), 3.76 (s, 2H), 3.24 (t, $J=4.7$ Hz, $2\times$ 2H), 2.72 (t, $J=$ 4.7 Hz, 2×2 H), 2.65 (s, 3H); ¹³C NMR (62.5 MHz, CDCl3): d 197.5, 157.6, 156.1, 151.1, 132.6, 129.0, 128.3, 124.7, 122.0, 119.7, 116.1, 111.2, 106.2, 55.3, 53.0, 49.0, 26.7; MS m/z (relative intensity) 334 (M⁺, 25), 228 (11), 201 (55), 173 (64), 130 (36), 106 (87), 56 (100). HRMS Calcd for $C_{21}H_{22}N_2O_2$ 334.1681, found 334.1690.

4.6.12. 1,4-Bis(benzofuran-2-ylmethyl)piperazine. Mp 143–144 °C; ¹H NMR (250 MHz, CDCl₃): δ 7.52–7.44 (m, 4H), 7.29–7.15 (m, 4H), 6.58 (s, 2×1 H), 3.69 (s, $2 \times$ 2H), 2.62 (s, br, 4×2 H); ¹³C NMR (62.5 MHz, CDCl₃): δ 155.0, 154.3, 128.2, 123.8, 122.6, 120.6, 111.2, 105.6, 55.3, 52.7; MS m/z (relative intensity) 346 (M⁺, 2), 215 (46), 131 (100), 77 (15). Anal. Calcd for $C_{22}H_{22}N_{2}O_{2}$: C, 76.28; H, 6.40; N, 8.09. Found: C, 76.05; H, 6.51; N, 7.99.

4.7. General procedure for recycling

After carried out a Mannich condensation, ether was added to remove the product from the cuprous iodide and alumina

surface. After filtration, the Cu/Al₂O₃ was directly used for the next trial.

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Chiral bisphospholane ligands (Me-ketalphos): synthesis of their Rh(I) complexes and applications in asymmetric hydrogenation

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Abstract—Rhodium complexes of functionalized bisphospholane ligands (S, S, S, S) -Me-ketalphos) 1 and (R, S, S, R) -Me-ketalphos) 2 have been used as catalyst precursors for the asymmetric hydrogenation of several different types of functionalized olefins and have achieved high enantioselectivities.

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1. Introduction

Development of new chiral phosphine ligands has played a significant role in the transition metal catalyzed asymmetric reactions during the last several decades.^{[1](#page-99-0)} Chiral C_2 -symmetric bisphospholane ligands such as $DuPhos²$ $DuPhos²$ $DuPhos²$ and its analogues RoPhos,^{[3a](#page-100-0)} BASPhos^{[3b,4](#page-100-0)} and MalPhos^{[5](#page-100-0)} have attracted much attention due to their effectiveness for asymmetric hydrogenation of functiona-lized olefins and ketones.^{[6](#page-100-0)} Recently, our group^{[7,8](#page-100-0)} and RajanBabu^{[9](#page-100-0)} reported a series of modified DuPhos ligands with ketal (1 and 2, named as ketalphos, prepared from inexpensive D-mannitol) or hydroxyl groups at the 3 and 4 positions of the phospholanes. In our previous reports, only one enantiomer of the ligands was synthesized. Designing a catalytic system that can obtain both enantiomers of the hydrogenation products is desired. Since preparation of the enantiomer of 1 may require the expensive L-mannitol, it is more practical to use its diastereomer 2 as ligand candidate to achieve opposite enantioselectivities in asymmetric hydrogenation. Rhodium complexes prepared in situ with ligand 1 and 2 gave good ee's in asymmetric hydrogenation reactions[.10](#page-100-0) However, only some preliminary results for the asymmetric hydrogenation of a-dyhydroamino acid derivatives were given. A systematic study of both diastereomers of Me-ketalphos in the hydrogenation of different kinds of unsaturated compounds is of significant importance.

According to the literature, $2c,11$ the use of in situ metal– ligand complexes generally leads to slightly less ee values in

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comparison to the corresponding isolated complexes. A possible reason is that the formation of the achiral rhodium catalyst species competes with the chiral active catalyst. In our previous attempts to prepare the complexes in situ, we found that the catalysts were inactive in the asymmetric hydrogenation possibly due to the short incubation time of metal precursor and the ligands. Based on a more recent report, $\frac{9}{5}$ we were able to obtain the active species by extending the incubation time. However, the asymmetric hydrogenation results through the in situ method of mixing metal precursors with ligands 1 and 2 leads to relatively lower ee's compared with the ones achieved by the isolated complexes. Also, the results of the in situ method were not easily reproducible. In this paper, we reported our studies based on the preparation of isolated metal complexes of Me-ketalphos ligands for asymmetric hydrogenation. The advantage of using the isolated complexes is high reproducibility, enantioselectivity and ease of operation (Fig. 1).

 $(2R, 3S, 4S, 5R)$ -Me-Ketalphos (2S, 3S, 4S, 5S)-Me-Ketalphos

Figure 1. Structure of both diastereomers of Me-ketalphos.

2. Results and discussion

(S,S,S,S)-Me-ketalphos and its diasteromer were prepared from commercially available D -mannitol.^{[8,9](#page-100-0)} Since the Rh-NBD complex precursor is more reactive than the Rh-COD

Keywords: Hydrogenation; Catalyst; Rhodium.

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$$
[Rh(NBD)_2]BF_4 + Ligand \xrightarrow{CH_3OH, rt} [Rh(NBD)(ligand)]BF_4
$$

3
4: ligand = 1
5: ligand = 2

Scheme 1. Synthesis of rhodium bisphospholane complex 4 and 5.

complex in the initiation reaction, 12 we selected $[Rh(NBD)_2]BF_4$ as the complex precursor in our catalyst precursor preparation (Scheme 1). To a solution of ligand in methanol was added 1 equiv of the complex precursor and stirred at rt for 15 min. After removal of the solvent, the Rh-Me-ketalphos complex gave a clean phosphorus NMR at 100 ppm. These Rh-Me-ketalphos complexes were characterized by ${}^{1}H$, ${}^{31}P$ NMR and HRMS.

Using an enamide 6a as the substrate, we have performed the condition optimization. As shown in Table 1, almost all the solvents gave good to excellent ee's except for ethanol (87% ee). In screening of solvents, we initially used methanol and found out that the results were not very consistent.[13](#page-100-0) Both methylene chloride and isopropanol were found to be solvents of the choice.

Table 1. Rhodium catalyzed asymmetric hydrogenation of an enamide 6a^a

^a The reactions were carried out at rt under 10 atm of H_2 pressure for 24 h with 100% conversions.

b Enantiomeric excesses were determined by chiral GC using a Supelco Chiral Select 1000 (0.25 mm \times 30 m) column.

Under the optimized conditions, we have investigated asymmetric hydrogenation of enamides with both 4 and 5 as catalysts (Table 2). Complete conversions and very high enantioselectivities were observed with complex 4 (92–98% ee's). The reaction time was not optimized to ensure the complete conversion. A more detailed study on substrate 6d showed that a TON of 1000 can be achieved without any deterioration of the enantioselectivity 14 and a TOF of $70 h^{-1.15}$ $70 h^{-1.15}$ $70 h^{-1.15}$ Catalyst 5 also behaved in a similar fashion, even though the enantioselectivities were slightly lower.

For hydrogenation of dehydroamino acid derivatives, complex 5 was found to be an excellent complex and high ee's for the substrates in Table 3 have been achieved (entries 1–7). In asymmetric hydrogenation, tetra-substituted substrates usually are more challenging in both reactivity and selectivity compared with tri-substituted ones. In our case, entry 8 showed high enantioselectivity (92%) under mild reaction condition, which is comparable to the best results

 a The reactions were carried out at rt under 10 atm of H_2 pressure for 24 h with 100% conversions.

^b Enantiomeric excesses were determined by chiral GC using a Supelco Chiral Select 1000 (0.25 mm \times 30 m) column.

 \degree The absolute configurations were assigned by comparision of the sign of optical rotation with reported data.

Table 3. Asymmetric hydrogenation of dehydroamino acid derivates by 4 and 5°

R^2 _COOCH_3 + H ₂ NHAc	4 or 5 (1 mol %) CH ₂ Cl ₂ , 12 h, rt	\mathcal{L} COOCH ₃ NHAc

^a The reactions were carried out at rt under 3 atm of H₂ pressure for 12 h. b Enantiomeric excesses were determined by chiral GC using a Chirasil-Val lll column.

^c The absolute configurations were assigned by comparing the sign of optical rotation with reported data.

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Table 4. Asymmetric hydrogenation of itaconic acid derivatives by complex 4 and 5^{a}

^a The reactions were carried out at rt under 10 atm of H₂ pressure for 24 h with 100% conversions. b Enantiomeric excesses were determined by chiral GC using a Gamma-DEX 225 column.

^c The absolute configurations were assigned by comparing the sign of optical rotation with reported data.

^d Enantiomeric excesses were determined on the corresponding methyl esters.

that have been reported.^{[7,16](#page-100-0)} However, the complex 4 of diastereomeric phospholane 1, only gave opposite selectivities in much lower ee values. The detailed reason for these results is still not clear. We have also explored hydrogenation of itaconic acid derivatives with catalysts 4 and 5. Excellent results (up to $> 98\%$ ee) were achieved with 10a and 10b (Table 4).

3. Conclusion

In summary, rhodium complexes 4 and 5 of (S, S, S, S) -Meketalphos and its diastereomer (R, S, S, R) -Me-ketalphos were prepared and characterized by NMR spectroscopy. The investigation of 4 and 5 in asymmetric hydrogenation of several different types of functionalized alkenes showed that catalyst 4 was highly enantioselective for the hydrogenation of enamides, while catalyst 5 performed very well for both enamides and dehydroamino acid derivatives.

4. Experimental

4.1. General methods

All reactions and manipulations were performed in a nitrogen-filled glovebox or using standard schlenk techniques. GC analysis was carried out using chiral capillary columns: Chirasil-Val III FOST (Dimensions: $30 \text{ m} \times$ 0.25 mm) for dehydroamino acid derivatives; Chiral Select 1000 column (dimensions: $15 \text{ m} \times 0.25 \text{ mm}$) for enamides; γ -225 (dimensions: 30 m \times 0.25 mm) for itaconic acid derivatives. Chemical shifts were reported in ppm downfield from tetramethylsilane with the solvent resonance as the internal standard.

4.2. General procedure for the synthesis of 4 and 5

To a solution of Me-ketalphos 1 (135 mg, 0.3 mmol) in 20 mL of methanol was added $(Rh(NBD)_2)BF_4$ (112.2 mg, 0.3 mmol) [NBD = norbornadiene], and the resulting bright orange solution was stirred at rt for 15 min. Solvent was removed under reduced pressure.

4.2.1. Rh(NBD)(1)]BF₄. ¹H NMR (360 MHz, CD₃OD) δ 7.91–7.96 (m, 2H), 7.78–7.82 (m, 2H), 6.13 (s, 2H), 5.91 (s, 2H), 4.72 (dd, $J=8.04$, 10.49 Hz, 2H), 4.29 (dd, $J=7.41$, 10.31 Hz, 4H), 3.26–3.32 (m, 2H), 2.77–2.86 (m, 2H), 1.93

(s, 2H), 1.55 (s, 6H), 1.54 (s, 6H), 1.30–1.37 (m, 6H), 0.73–0.79 (m, 6H). ³¹P NMR (CD₃OD) δ 100.1 (d, $J_{\text{Rh-P}}$ 153.5 Hz); HRMS (cation) m/z calcd for $C_{31}H_{44}O_4P_2Rh$ 645.1770, found 645.1737; HRMS (anion) m/z calcd for BF₄ 87.0029, found 87.0024.

4.2.2. Rh(NBD)(2)]BF₄. ¹H NMR (360 MHz, CD₃OD) δ 8.04–8.28 (m, 2H), 7.86–7.88 (m, 2H), 6.14 (s, 2H), 4.40 (s, 2H), 4.28 (dd, $J=10.1$, 18.9 Hz, 2H), 3.76 (dd, $J=8.9$, 11.4 Hz, 4H), 2.81–2.92 (m, 2H), 2.62–2.79 (m, 2H), 2.06 (s, 2H), 1.64 (s, 6H), 1.62 (s, 6H), 1.30–1.60 (m, 6H), 0.87–0.93 (m, 6H). ³¹P NMR (CD₃OD) δ 100.8 (d, $J_{\text{Rh-P}}=$ 158.0 Hz); HRMS (cation) m/z calcd for $C_{31}H_{44}O_4P_2Rh$ 645.1770, found 645.1752; HRMS (anion) m/z calcd for BF_4 87.0029, found 87.0021.

4.3. General hydrogenation procedure using 4 or 5

In a glove box, a solution of 0.005 mmol of catalyst in 1 mL of CH_2Cl_2 was added 0.5 mmol of substrate. The resulting mixture was transferred to an autoclave. The hydrogenation was performed at rt under 3–10 atm of hydrogen for 12–24 h. The hydrogen was carefully released and the reaction mixture was passed through a short silica gel plug to remove the catalyst. The ee were measured by GC with a chiral column directly without any further purification. The absolute configurations of the products were determined by comparing the sign of optical rotation with the reported values.

All the physical characterization data for substrates and products can be found in Ref. [17](#page-100-0) and the references therein.

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- 13. Phosphorus NMR showed that the complex was decomposed after certain period of time with detection of a small peak at 83 ppm.
- 14. The hydrogenation reaction was performed with 0.1 mol% loading of complex 4 at standard reaction condition for enamides for 16 h with 100% conversion.
- 15. The hydrogenation reaction was performed with 0.1 mol% loading of complex 4 at standard reaction condition for enamides for 2 h with 14% conversion.
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A highly active ionic liquid catalyst for Morita–Baylis–Hillman reaction

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Abstract—The Morita–Baylis–Hillman reaction is an efficient carbon–carbon bond forming reaction for the preparation of α -methyleneb-hydroxycarbonyl compounds. A new and highly active di-naphthalene imidazolium salt has been synthesized. We have found that 1,3-bis[2-(naphthalene-2-yloxy)propyl]imidazolium bromid promoted the Morita–Baylis–Hillman reaction of various aryl aldehyde compounds in the absence of solvents. Our studies show that the Morita–Baylis–Hillman reaction by the influence of ionic liquid to give a high yield and short reaction time.

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1. Introduction

The Morita–Baylis–Hillman (MBH) reaction is a particularly powerful transformation because it permits the coupling of various activated aldehydes and α , β -unsaturated ketones, a process that has a great synthetic advantage.^{[1](#page-105-0)} This reaction involves three components: an activated alkene, an electrophile, and a tertiary amine. Lewis bases, such as 1,[4](#page-105-0)-diazabicyclo[2.2.2]octane $(DABCO)$, 2 DMAP, 3 DBU, 4 $2,2'$ -bis(diphenylphosphino)-1,1'-binaphthyl (BINAP),^{[5](#page-105-0)} and $imidazole⁶$ $imidazole⁶$ $imidazole⁶$ are frequently used in these reactions as catalysts. In addition, Lewis acid-accelerated reactions were also reported, such as $TiCl₄$, 7 7 Et₂AlI,^{[8](#page-106-0)} and BF₃.^{[9](#page-106-0)} In a general process, the MBH reaction suffers from poor reaction rates and long reaction time. For this reason, modifications such as the use of high pressure,^{[10](#page-106-0)} microwave irradiation^{[11](#page-106-0)} and ultrasound^{[12](#page-106-0)} have given some promising results. In recent years, room temperature ionic liquids (RTILs) have been gaining exposure for their potential in organic synthesis. From both an environmental and an economical point of view, the concept of recoverable and recyclable catalysis has become increasingly important.^{[13](#page-106-0)} RTILs have received considerable attention for applications as green solvents^{[14](#page-106-0)} and replacements for traditional organic

A common MBH reaction carried out with [bmim][X] to give lower yield $(13-78\%)^{26b}$ $(13-78\%)^{26b}$ $(13-78\%)^{26b}$ and a longer reaction time.

$$
- \frac{N}{N} \left(\frac{1}{N} \right)_{N} R \left[\text{CI} \right]
$$

R = CO₂H, SOCl

Scheme 1. Ionic liquids.

Keywords: Morita–Baylis–Hillman reaction; Ionic liquids; DABCO.

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Table 1. Reaction of various aldehydes with methyl vinyl ketone in the presence of [bmim][Br] or [bmim][BF4]

We prepared [bmim][$Br]^{28a}$ $Br]^{28a}$ $Br]^{28a}$ and [bmim][BF_4]^{[28b](#page-106-0)} and then carried out the reactions of these RTILs with aldehydes and α , β -unsaturated ketones in the presence of DABCO (Table 1). Since we were not satisfied with the results, we

synthesized a new imidazolium salt for the MBH reaction (Scheme 2). The reaction of methyl lactate with 2-naphthol in the presence of DEAD^{[29](#page-106-0)} is followed by sodium borohydride reduction yields the primary alcohol 1.^{[30](#page-106-0)}

Scheme 2. Synthesis of di-naphthalene imidazolium salt.

Table 2. Reaction of various aldehydes with methyl vinyl ketone in the presence of 1,3-bis[2-(naphthalene-2-yloxy)propyl]imidazolium bromide

$$
R \n\begin{matrix}\n0 & 0 & 4, DABCO \\
H + \n\end{matrix}\n\begin{matrix}\n0 & 0 & 0 \\
1, t & 0\n\end{matrix}
$$

OH \overline{O}

Table 3. Reaction of various aldehydes with methyl acrylate in the presence of 1,3-bis[2-(naphthalene-2-yloxy)propyl]imidazolium bromide

 \overline{O}

The tosylation of the primary alcohol followed by a treatment with 1.3 equiv of lithium bromide in acetone produced bromide 2. The reaction of imidazole with compound 2 afforded the imidazole derivative, isolated as compound 3. Finally, the treatment of 3 with bromide 2 at 60° C gave $1,3$ -bis[2-(naphthal-2-yl-oxy)propyl]imidazolium bromide 4 in 96% yield after purification.

 $\Omega_{\rm H}$

The treatment of various aryl aldehydes with methyl vinyl ketone (MVK) or methyl acrylate in the presence of hydrophilic ionic liquid 4 not only produced the desired product in moderate to high yield but also reduced the side products. For example, in the reaction of methyl 4-formylbenzoate (1.0 equiv) with MVK (2.0 equiv), entry 2 ([Table 2](#page-102-0)) can be obtained in 96% yield in the presence of compound 4 (0.2 equiv) for 0.5 h at room temperature. We also prepared MBH reaction with methyl acrylate (Table 3), our condition goes very smoothly, producing high yields and also accelerating the reaction rate. The recovered ionic liquid was reused for subsequent runs.^{[16b](#page-106-0)}

In conclusion, the present procedure catalyzed by a hydrophilic RTIL, 1,3-bis[2-(naphthal-2-yl-oxy)propyl] imidazolium bromide provides an efficient and general methodology for the MBH reaction of aryl aldehydes and MVK or methyl acrylate to yield α -methylene- β -hydroxycarbonyl compounds. The results presented in this paper, demonstrate that the MBH reaction works well in the presence of ionic liquid.

2. Experimental

All reagents were purchased from Acros, Aldrich Chimie, Fluka and used without further purification. IR spectra were taken as KBr plates. ${}^{1}H$ and ${}^{13}C$ NMR spectra were recorded in CDCl₃ solution at 300 and 75 MHz spectrometer. TLC analysis was performed used glass plate with silica gel 60 $F₂₅₄$. Flash column chromatography was carried out using 200–300 mesh silica gel.

2.1. Typical procedure for the ionic liquid promoted Morita–Baylis–Hillman reaction

In a dried flask, aldehyde (1 mmol), DABCO (224 mg, 2 mmol), and 1,3-bis[2-(naphthal-2-yl-oxy)propyl]imidazolium bromide (129.3 mg, 0.25 mmol) were measured together. Methyl vinyl ketone (105 mg, 1.5 mmol) was added, and the reaction mixture was stirred for 7–20 h at ambient temperature in all cases. The mixture was washed with ether $(3 \times 10 \text{ mL})$, and the solid was separated by vacuum filtration. Then the solid was extracted with 130 mL dichloromethane and followed by a general procedure to recover the catalyst. The ethereal phase was concentrated, and the solvent was removed under reduced pressure. The residue was purified by silica gel column chromatography (Hexane/ethyl acetate, 7:3, v/v).

2.1.1. Synthesis of 1. Methyl lactate $(10.40 \text{ g}, 0.1 \text{ mol})$ was added to a solution of 2-naphthol (14.40 g, 0.11 mmol) and triphenyl phosphine (31.44 g, 0.12 mol) in dichloromethane (100 mL). The reaction mixture was stirred at 0° C and diethyl azodicarboxylate (17.04 g, 0.12 mol) was added. After the mixture was stirred for 2 h at room temperature and the white solid ($Ph_3P=O$) was separated by vacuum filtration. The organic layer was evaporated under vacuum, the crude dissolved in a co-solvent (200 mL tetrahydrofuran and 20 mL methol), sodium tetrahydridoborate (3.8 g, 0.1 mol) was added, and stirred at ambient temperature for 3.5 h. The organic phase was washed with 100 mL of water and dried over anhydrous MgSO₄. The organic solvent was evaporated under reduced pressure to give 2-(1-methoxypropan-2-yloxy)naphthalene (17.56 g, 81%) as yellow oil.

IR (KBr) v_{max} : 3420 (OH), 3056, 1215, 2975, 2931, 1627, 1598, 1509, 1467, 1256, 1215, 1180, 1050, 970, 839, 748; ¹H NMR (300 MHz, CDCl₃) δ : 1.39–1.37 (d, J=7.0 Hz, $3H$, $3.85-3.82$ (d, $J=7.8$ Hz, 2H), 4.70–4.63 (m, 1H), 7.22–7.15 (m, 2H), 7.45–7.35 (m, 2H), 7.79–7.71 (m, 2H); ¹³C NMR (CDCl₃) δ : 155.59, 134.60, 129.71, 127.74, 126.88, 126.53, 123.93, 119.62, 109.03, 74.83, 66.20, 15.81; MS (EI, 70 eV, m/z , %): 202 (M⁺, 65), 171 (28), 143 (100),

127 (55), 115 (77), 101 (11); HRMS calcd for $C_{13}H_{14}O_2$: 202.0994. Found: 202.0998.

2.1.2. Synthesis of 2. A mixture of 2-(1-methoxypropan-2-yloxy)naphthalene (10.80 g, 0.05 mol), tosyl chloride (11.44 g, 0.06 mol), and pyridine (5.3 mL, 0.06 mol) was stirred in 100 mL of dichloromethane at room temperature for 18 h. The organic phase was washed with 150 mL water, dried over anhydrous MgSO₄ and concentrated. Then the tosylate crude mixture wit lithium bromide (5.66 g, 0.065 mol) was diluted in 50 mL of acetone, and lithium carbonate (0.37 g, 0.05 mol) added. The mixture was stirred at ambient temperature for 24 h. When the reaction was completed by a TLC analysis, the solvent was evaporated and a dark yellow oil was purified via flash chromatography (Hexane/ethyl acetate, 9:1, v/v), producing a light yellow oil, 2-(1-bromopropan-2-yloxy)naphthalene (12.28 g, 93%).

IR (KBr) ν_{max} : 1627, 1598, 1509, 1466, 1255, 1214, 1177, 838, 747; ^TH NMR (300 MHz, CDCl₃) δ : 1.54 (d, $J=6.2$ Hz, 3H), 3.53–3.48 (m, 1H), 3.67–3.61 (m, 1H), 4.76–4.70 (m, 1H), 7.19–7.15 (m, 2H), 7.39–7.36 (m, 1H), 7.46–7.43 (m, 1H), 7.79–7.72 (m, 3H); ¹³C NMR (CDCl₃) d: 155.12, 134.43, 129.77, 129.13, 127.67, 126.82, 126.49, 124.02, 119.54, 109.15, 73.53, 35.31, 18.79; MS (FAB, m/ z): 266 (M^+) , 264, 145, 144, 127; HRMS calcd for $C_{13}H_{13}BrO: 264.0150.$ Found: 264.0151.

2.1.3. Synthesis of 3 and 4. Bromide 2 (5.30 g, 0.02 mol) was injected into a flask containing tetrahydrofuran (55 mL) followed by potassium carbonate (0.55 g, 8 mmol) and stirred at room temperature for 24 h. After completion of the reaction, as indicated by TLC, the product was quenched with 50 mL water and extracted with dichloromethane ($3 \times$ 50 mL). The organic layer was dried over anhydrous MgSO4, concentrated under reduced pressure and purified by chromatography on silica gel to afford imidazole 3 (4.59 g, 91%). The combined bromide 2 (2.52 g, 0.01 mol) and imidazole 3 (2.65 g, 0.01 mol) was stirred at 60 °C for 2 h. The mixture was washed with diethyl ether $(3 \times$ 100 mL) and evaporated under vacuum. A brown solid, 1,3 bis[2-(naphthalene-2-yloxy)propyl]imidazolium bromide, was obtained (9.93 g, 96%).

Compound 3. IR (KBr) v_{max} : 2924, 1735, 1508, 1255, 1214, 748; ¹H NMR (CDCl₃) δ : 1.37 (d, J=6.0 Hz, 3H), 4.22–4.19 (m, 2H), 4.70–4.80 (m, 1H), 7.09–7.03 (m, 4H), 7.44–7.35 (m, 2H), 7.58 (s, 1H), 7.78–7.68 (m, 3H); 13C NMR (CDCl3) d: 156.52, 137.89, 134.00, 129.89, 128.62, 129.60, 127.69, 126.75, 126.58, 124.11, 120.00, 119.31, 108.95, 73.21, 51.66, 17.15; MS (EI, 70 eV, m/z, %): 252 $(M⁺, 83)$, 171 (46), 144 (93), 127 (56), 109 (100); HRMS calcd for $C_{16}H_{16}N_2O$: 252.1263. Found: 252.1259.

Compound 4. IR (KBr) v_{max} : 3056, 2195, 1626, 1598, 1509, 1467, 1255, 1180, 1119, 1078, 972, 732; ¹H NMR (CDCl₃) δ : 1.44 (d, J = 6.2 Hz, 6H), 4.47–4.39 (m, 2H), 4.96–4.89 (m, 4H), 7.01–6.98 (m, 2H), 7.13–7.12 (m, 2H), 7.44–7.34 (m, 8H), 7.71–7.62 (m, 8H); ¹³C NMR (CDCl₃) δ : 154.15, 132.24, 129.92, 128.80, 127.60, 126.88, 126.68, 124.27, 122.56, 118.80, 109.13, 72.41, 67.99, 54.21, 25.62, 16.63; MS (FAB, m/z): 517 (M)⁺, 437 (M-Br)⁺, 293, 253, 185,

144, 109. Anal. Calcd for $C_{29}H_{29}BrN_2O_2$: C, 67.31; H, 5.65. Found: C, 67.35; H, 5.68; mp: 178 °C.

Compounds 5^{31} 5^{31} 5^{31} , 7^{31} , 9^{32} 9^{32} 9^{32} , 11^{33} 11^{33} 11^{33} , 12 – 13^{34} 13^{34} 13^{34} , 16 – 17^{35} 17^{35} 17^{35} , 20^{36} 20^{36} 20^{36} , 22^{34} 23,^{[37](#page-106-0)} 24,^{[38](#page-106-0)} 26,^{[12a](#page-106-0)} 27,^{[39](#page-106-0)} 28–29,^{[40](#page-106-0)} and 31–32,^{[41](#page-106-0)} were identical to authentic samples by comparison of their spectral date.

2.1.4. Methyl 4-[(1-hydroxy)(2-methylene)butyl-3-one] **benzoate** (6). IR (KBr) ν_{max} : 3460 (OH), 1719 (C=O), 1677 (C=O), 1434, 1283, 1115, 960, 766, 707; ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$ δ : 8.01 (d, J=6.6 Hz, 2H), 7.44 (d, $J=8.3$ Hz, 1H), 6.22 (s, 1H), 5.97 (s, $J=5.97$ Hz, 1H), 5.65 $(s, 1H), 3.91 (s, 1H), 2.34 (s, 1H);$ ¹³C NMR (75.4 MHz, CDCl₃) δ : 200.29 (C=O), 168.89, 149.46, 146.63, 129.72, 129.48, 127.32, 126.44, 72.64, 52.13, 26.44; MS (FAB, m/z): 217 ($M^+ + 1 - H_2O$); HRMS calcd for $C_{13}H_{14}O_4$: 234.0892. Found: 234.0897.

2.1.5. 3-(Hydroxy(naphthalene-2-yl)methyl)but-3-en-2 one (8). IR (KBr) ν_{max} : 3436 (OH), 3055, 1672 (C=O), 1363, 1125, 1041, 952, 824, 747; ¹H NMR (300 MHz, CDCl₃) δ : 7.85–7.80 (m, 4H), 7.49–7.23 (m, 4H), 6.23 (s, 1H), 6.01 (s, $J=1.1$ Hz, 1H), 5.80 (s, 1H), 2.35 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ: 200.46 (C=O), 149.92, 138.84, 133.26, 132.92, 128.17, 128.12, 127.67, 127.00, 126.18, 126.03, 125.47, 124.56, 72.91, 26.54; MS (EI, 70 eV, m/z , %): 226 (M⁺, 84), 225 (100), 211 (14), 166 (15), 165 (34), 155 (23), 128 (54), 127 (43), 84 (50); HRMS calcd for $C_{17}H_{16}O_2$: 226.0994. Found: 226.0996.

2.1.6. 3-[Hydroxy(4-phenylphenyl)methyl]but-3-en-2 one (10). IR (KBr) ν_{max} : 3436 (OH), 3029, 1672 (C=O), 1485, 1363, 1038, 763, 700; ¹H NMR (300 MHz, CDCl₃) δ : 7.60–7.56 (m, 4H), 7.45–7.43 (m, 4H), 7.42 (d, 1H), 6.23 (s, 1H), 6.05 (d, $J=1.1$ Hz, 1H), 2.37 (s, 3H); ¹³C NMR $(75.4 \text{ MHz}, \text{CDCl}_3)$ δ : 200.39 (C=O), 149.90, 140.80, 140.65, 140.59, 128.78, 127.33, 127.12, 127.11, 126.97, 126.80, 72.73, 26.55; MS (EI, 70 eV, m/z , %): 252 (M⁺ 87), 251 (100), 191 (22), 175 (75), 152 (35), 77 (26); HRMS calcd for $C_{17}H_{16}O_2$: 252.1150. Found: 252.1150.

2.1.7. 2-[Hydroxy(3,4,5-trimethoxyphenyl)methyl] cyclohex-2-enone (14). IR (KBr) v_{max} : 3480 (OH), 2937, 2838, 1672 (C=O), 1591, 1506, 1459, 1418, 1325, 1231, 1125, 1008, 843; ¹H NMR (300 MHz, CDCl₃), δ 6.58 (s, 2H), 6.17 (s, 1H), 5.96 (s, 1H), 3.85, (s, 6H), 3.83 (s, 3H), 2.36 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 200.53 (C=O), 153.24, 149.90, 137.36, 137.10, 126.88, 103.54, 72.75, 60.83, 56.12, 26.56; MS (70 eV, m/z , %): 266 (M⁺ 54), 235 (74), 196 (100), 181 (49), 169 (28), 138 (18), 125 (31), 110 (19), 84 (70); HRMS calcd for $C_{14}H_{18}O_5$: 266.1154. Found: 266.1147.

2.1.8. 3-[Hydroxy(4-methylthiophenyl)methyl]but-3-en-**2-one (15).** IR (KBr) ν_{max} : 3437 (OH), 2920, 1673 (C=O), 1596, 1493, 1403, 1364, 1092, 1038, 972, 820; ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$ δ : 7.29–7.20 (dd, 4H), 6.19 (s, 1H), 5.99 (s, 1H), 5.58 (s, 1H), 2.49 (s, 3H), 2.34 (s, 3H); 13C NMR $(75.4 \text{ MHz}, \text{CDCl}_3)$ δ : 200.33 (C=O), 149.88, 138.46, 137.87, 127.08, 126.63, 126.57, 72.46, 26.52, 15.84; MS (EI, 70 eV, m/z , %): 222 (M⁺, 39), 221 (42), 207 (8), 175 (100), 153 (37), 151 (37), 124 (21), 109 (30), 77 (18); HRMS calcd for $C_{12}H_{14}O_2S$: 222.0715. Found: 222.0718.

2.1.9. 3-[(4-Bromothiophen-2-yl)(hydroxy)methyl]but-3 en-2-one (18). IR (KBr) v_{max} : 3421 (OH), 3106, 2919, 1672 $(C=0)$, 1367, 1033, 971, 821; ¹H NMR (300 MHz, CDCl₃) δ : 7.16 (s, J = 1.4 Hz, 1H), 6.85 (s, J = 1.2 Hz, 1H), 6.38 (s, 1H), 5.96 (s, 1H), 5.68 (s, 1H), 3.77 (s, 3H); 13C NMR $(75.4 \text{ MHz}, \text{CDCl}_3)$ δ : 166.37 (C=O), 147.28, 140.35, 127.14, 127.03, 122.59, 109.35, 69.83, 52.27; MS (FAB, m/z): 243 ($M^+ + 1 - H_2O$); HRMS calcd for C₉H₉BrO₂S: 259.9507. Found: 259.9512.

2.1.10. 3-{[5-(3-Chlorophenyl)furan-2-yl](hydroxy) methyl}but-3-en-2-one (19). IR (KBr) v_{max} : 3429 (OH), 2919, 2365, 1672 (C=O), 1605, 1430, 1367, 1019, 781; ¹H NMR (300 MHz, CDCl₃) δ : 7.6 (d, J = 1.3 Hz, 1H), 7.48 (t, 1H), 7.31–7.19 (m, 2H), 6.61 (d, $J=3.3$ Hz, 1H), 6.34 (d, $J=3.3$ Hz, 1H), 6.29 (s, 1H), 6.17 (s, 1H), 5.67 (s, 1H), 2.41 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 200.02 (C=O), 154.48, 152.17, 147.05, 134.69, 132.28, 129.96, 127.70, 123.70, 127.33, 121.80, 109.44, 106.94, 67.51, 26.42; MS $(70 \text{ eV}, \text{m/z}, \%)$: 278 $(M+2, 33)$, 276 $(M^+, 100)$, 247 (6) , 233 (19), 215 (18), 207 (72), 191 (12), 178 (74), 165 (8), 152 (21), 149 (32), 139 (89), 137 (40), 115 (69), 111 (29), 98 (81), 70 (38); HRMS calcd for $C_{15}H_{13}ClO_3$: 276.0553. Found: 276.0556.

2.1.11. Methyl 4-[2-(methoxycarbonyl)hydroxyallyl] **benzoate (21).** IR (KBr) v_{max} : 3491 (OH), 2955, 1723 $(C=0)$, 1436, 1283, 1110, 1019, 818; ¹H NMR (300 MHz, CDCl₃) δ : 8.03 (s, 1H), 8.00 (s, 1H), 7.47 (s, 1H), 7.45 (s, 1H), 6.36 (s, 1H), 5.83 (s, 1H), 5.60 (s, 1H), 3.91 (s, 1H), 3.71 (s, 1H); ¹³C NMR (75 MHz, CDCl₃) δ : 166.9 (C=O), 166.6 (C=O), 146.3, 141.4, 129.8, 129.6, 126.9, 126.5, 73.13, 52.2, 52.1; MS (EI, 70 eV, m/z , %): 250 (M⁺, 35), 235 (18), 219 (25), 191 (31), 190 (35), 165 (18), 163 (100), 131 (13), 105 (31), 77 (30), 59 (21); HRMS calcd for $C_{13}H_{14}O_5$: 250.0841. Found: 250.0849.

2.1.12. Methyl 2-[hydroxy(4-phenylphenyl)methyl] acrylate (25). IR (KBr) v_{max} : 3370 (OH), 2952, 1719 $(C=0)$, 1434, 1268, 1151, 1030, 766; ¹H NMR (300 MHz, CDCl₃) δ : 7.64 (m, 4H), 7.50 (m, 4H), 7.37 (t, J=1.2 Hz, 1H), 6.38 (s, 1H), 5.91 (s, 1H), 5.62 (s, 1H), 3.75 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 166.82 (C=O), 141.84, 140.78, 140.76, 140.30, 128.80, 127.37, 127.25, 127.12, 127.03, 126.30, 73.18, 52.06; MS (EI, 70 eV, m/z, %): 268 $(M⁺, 70)$, 236 (22), 208 (21), 191 (27), 181 (100), 153 (20), 77 (18); HRMS calcd for $C_{17}H_{16}O_3$: 268.1099. Found: 268.1097.

2.1.13. Methyl 2-{hydroxy[4-(methylthio)phenyl] methyl}acrylate (30). IR (KBr) ν_{max} : 3458 (OH), 2952, 2919, 1715 (C]O), 1631, 1495, 1434, 1287, 1435, 1287, 1146, 1037, 957, 818; ¹H NMR (300 MHz, CDCl₃) δ: 7.28 (dd, 2H), 7.22 (dd, 2H), 6.33 (s, 1H), 5.85 (d, $J=1.0$ Hz, 1H), 5.52 (s, 1H), 3.72 (s, 3H), 3.07 (d, 1H), 2.47 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 166.77 (C=O), 141.76, 138.07, 127.15, 126.45, 126.18, 72.92, 52.08, 15.75; MS (FAB, m/z): 221 (M⁺+1-H₂O); HRMS calcd for $C_{12}H_{14}O_3S: 238.0664.$ Found: 238.0668.

2.1.14. Methyl 2-[hydroxy(4-bromothiophen-2 yl)methyl]acrylate (33). IR (KBr) v_{max} : 3447 (OH), 3110, $2952, 1712$ (C=O), 1437, 1335, 1151, 1038,818; ¹H NMR

(300 MHz, CDCl₃) δ : 7.13 (s, J = 1.3 Hz, 1H), 6.83 (s, J = 0.9 Hz, 1H), 6.24 (s, 1H), 6.12 (s, 1H), 5.72 (s, 1H), 2.37 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 200.17 (C=O), 148.30, 147.49, 127.40, 127.04, 122.36, 109.30, 69.53, 26.50; MS (EI, 70 eV, m/z , %): 278 (M+2, 100), 276 (M⁺, 97), 261 (71), 246 (52), 218 (42), 191 (57), 166 (30), 121 (26), 109 (25), 97 (10), 84 (48); HRMS calcd for C9H9BrO3S: 275.9456. Found: 275.9453.

2.1.15. Methyl 2-[(5-(3-chlorophenyl)furan-2-yl)- (hydroxy)methyl]acrylate (34). IR (KBr) v_{max} : 3449 (OH), 2947, 1715 (C=O), 1628, 1438, 1286, 1202, 783; ¹H NMR (300 MHz, CDCl₃) δ : 7.6 (d, J = 1.7 Hz, 1H), 7.48 $(t, 1H), 7.31-7.19$ (m, 2H), 6.61 (d, $J=3.3$ Hz, 1H), 6.34 (d, $J=3.3$ Hz, 1H), 6.29 (s, 1H), 6.17 (s, 1H), 5.67 (s, 1H), 2.41 (s, 3H); ¹³C NMR (75.4 MHz, CDCl₃) δ : 200.02 (C=O), 154.48, 152.17, 147.05, 134.69, 132.28, 129.96, 127.70, 123.70, 127.33, 121.80, 109.44, 106.94, 67.51, 26.42; MS $(70 \text{ eV}, \frac{m}{z}, \frac{\%}{z})$: 294 $(M+2, 21)$, 292 $(M^+, 70)$, 260 (25) , 232 (27), 207 (89), 205 (70), 178 (74), 153 (40), 149 (40), 141 (77), 139 (100), 115 (78), 113 (30), 111 (31), 75 (22); HRMS calcd for $C_{15}H_{13}ClO_3$: 292.0502. Found: 292.0509.

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Complementary regioselective esterification of non-reducing oligosaccharides catalyzed by different hydrolases

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Abstract—The enzymatic transesterification of several tri- and tetrasaccharides with vinyl laurate is described. The lipases from Candida antarctica B (Novozym 435) and Thermomyces lanuginosus (Lipozyme TL IM) and the alkaline protease from Bacillus licheniformis (subtilisin Carlsberg) have been used with each carbohydrate to obtain different regioisomers. By using the sugars in their amorphous form, complete solubility is achieved in the reaction media (tert-butanol/pyridine mixtures for the lipases and pyridine for the protease) and high isolated yields of the corresponding monoesters are obtained. Good to excellent regioselectivity is observed for all the enzymes, showing a final complementary picture respect to the primary hydroxyls of the oligosaccharides studied.

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1. Introduction

Carbohydrate fatty acid esters are non-ionic surfactants with broad applications in food, cosmetic and pharmaceutical industries.^{[1](#page-114-0)} They are biodegradable, non-toxic and can be produced from inexpensive and renewable raw materials.^{[2](#page-114-0)} Their emulsifying and surfactant properties^{[3](#page-115-0)} may be modulated by the type of fatty acid, the sugar moiety, the degree of substitution and the position of attachment to the fatty acid (for instance regioisomeric sucrose monoesters present different CMC values).[4](#page-115-0)

Regioselective chemical acylation of carbohydrates is a particular challenging task due to their multifunctionality.[5](#page-115-0) For this reason, tedious multi-step synthesis based on protection/deprotection reactions is usually required.[6](#page-115-0) The use of enzymes in organic solvents has helped to solve this problem and both, lipases^{[7](#page-115-0)} and proteases^{[8](#page-115-0)} have been employed for the regioselective acylation of carbohydrates.^{[9](#page-115-0)} Different factors affect the activity, stability and selectivity of hydrolases in non-aqueous media.^{[10](#page-115-0)} The nature of the solvent is particularly important since a compromise between enzyme activity and saccharide solubility is required. This is even more important in the case of lipases that are active in hydrophobic solvents limiting acylation mainly to mono- and disaccharides. 11 To our knowledge, only three previous examples of enzymatic acylation of triand tetrasaccharides have been described in the literature. Riva et al.^{[12,13](#page-115-0)} synthesized a series of esterified tri- and tetrasaccharides by using subtilisin in dimethylformamide and trihaloethyl butyrates as the acylating agents to get ca. 29–50% isolated yields. Subtilisin showed a strong preference towards the regioselective esterification of the primary hydroxyls of the fructose unit in fructose-containing oligosaccharides and to the $6''$ -OH in the non-reducing glucose of maltotriose. Ferrer et al.^{[14](#page-115-0)} prepared long-chain fatty acid esters of maltotriose using the lipase from Thermomyces lanuginosus (previously Humicola Lanuginosa) in a medium constituted by two miscible solvents (2-methyl-2-butanol/dimethylsulfoxide). The hydroxyl 6 ⁿ-OH in the non-reducing end of the trisaccharide was acylated in low yields with 20% DMSO mixtures. Higher yields could be obtained by using 5% DMSO mixtures, but 10% of diesters were formed under these reaction conditions. Recently, Gustavsson et al. 15 reported similar reaction conditions using the lipase from Candida antarctica B for the transesterification of xyloglucan oligosaccharides mixtures with vinyl stearate obtaining 14% yield and some disubstitution (less than 10% of the total acylated products).

We were interested in preparing fatty acid esters of tri- and tetrasaccharides in order to study the relevance of the specific carbohydrate, its size and acylation position in the physicochemical properties of this type of non-ionic

Keywords: Oligosaccharide acylation; subtilisin Carslberg; Lipases; Candida antarctica; Thermomyces lanuginosus; Transesterification; Regioselectivity.

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 $2 : R = R' = R'' = H$ (melezitose) **2a**: $R = CO(CH_2)_{10}CH_3$; $R = R' = H$ **2b**: R = H ; R = $CO(CH_2)_{10}CH_3$; R" = H **2c**: R = R = H ; R = $CO(OH_2)_{10}OH_3$

4a: R = $CO(CH_2)_{10}CH_3$: R = H **4b**: R = H; R = $CO(CH_2)_{10}CH_3$

 $3 : R = R = R' = H (1 - 1)$ **3a**: R = CO(CH₂)₁₀CH₃; R' = R'' = H **3b**: R = H; R = $CO(CH_2)_{10}CH_3$; R" = H **3c**: R = R' = H; R" = $CO(OH_2)_{10}OH_3$

Scheme 1.

surfactants. Moreover, recently reported antibacterial activity of 6^{''}-O-laourylmaltotriose^{[16](#page-115-0)} and antitumoral activity of $6''$ -O-palmitoylmaltotriose^{[17](#page-115-0)} confirm the interest in developing selective methods for the preparation of large-oligosaccharide fatty acid esters. In this work, we present the enzymatic transesterification of the non-reducing oligosaccharides raffinose, melezitose, 1-kestose and stachyose with vinyl laurate. The reactions were performed in tert-butanol/pyridine mixtures with lipases from Thermomices lanuginosus (Lipozyme TL IM) and from Candida antarctica B (Novozym 435), and in pyridine with the alkaline protease from Bacillus licheniformis (subtilisin Carlsberg). The three enzymes studied gave high isolated yields of the sugar monoesters and only small amounts of diesters were produced. Regioselectivity for each oligosaccharide and enzyme is also discussed showing a complementary behaviour that allows the access to a particular regioisomer by selecting the appropriate hydrolase.

2. Results and discussion

All the commercially available non-reducing oligosaccharides considered in this investigation contain a sucrose moiety in their chemical structure (Scheme 1) The tri- and tetrasaccharides raffinose (1) and stachyose (4) can be considered as sucrose molecules substituted at the C-6 hydroxyl group while the trisaccharides melezitose (2) and

1-kestose (3) can be considered as a $C-3'$ hydroxylglucosylated sucrose and a $C-1'$ hydroxyl-fructosylated sucrose, respectively. This common structural feature led us to select the three different hydrolases indicated in the introduction because they have previously shown different

regioselectivity in the acylation of sucrose.[12,18](#page-115-0)

In all these enzyme-catalyzed transesterifications, vinyl laurate was chosen as acyl donor. We decided to use the amorphous form of each oligosaccharide obtained by lyophilization to increase the solubility of the sugars in organic solvents with the additional advantage of removing almost all the water of crystallization.^{[19](#page-115-0)} The regioisomeric distribution in the isolated products was determined by HPLC/MS (see Supplementary data) and the structure of main regioisomers was established by NMR. The position of esterification was determined by correlation bidimensional analysis ${}^{1}H-{}^{1}H$ (COSY) and ${}^{1}H-{}^{13}C$ NMR (HSQC, HMBC) and in some cases using 1D selective TOCSY and HSQC-TOCSY experiments.

Transesterifications catalyzed by Candida antarctica lipase B

First, we investigated the esterification of raffinose (1) with lipase Novozym 435, an immobilized preparation of C. antarctica lipase B on a macroporous acrylic resin. Due to the low solubility of trisaccharides in the relatively

apolar solvents needed to use lipases, we tried a tertbutanol/pyridine (55:45) mixture at 60 \degree C to carry out the enzymatic reaction. The specific ratio of solvents and temperature were selected since Novozym 435 shows the highest enzyme activity in the acylation of glucose with myristic acid in the above reaction conditions.^{[20](#page-115-0)}

Using 0.04 M of 1 and 0.2 M acyl donor as starting concentrations, complete solubilization of the sugar was observed in the reaction media, and 48% isolated yield of raffinose monoester was obtained after 72 h of reaction (Table 1). A small proportion of starting material and diester was also observed by TLC. Analysis of raffinose monolaurate showed that only the regioisomer possessing the ester substitution at the 6-OH of the galactose unit was formed. The reaction processing is very simple since it only requires filtration, rotary-evaporation of the solvents and column chromatography of the residue. Interestingly, only one regioisomer was obtained, whereas for sucrose this lipase produces a 1:1 mixture of the 6 and $6'$ esters.^{[11](#page-115-0)}

We extended the same reaction conditions to the other two trisaccharides: melezitose (2) and 1-kestose (3). Similarly to raffinose, a small proportion of diester and remaining starting sugar was detected by TLC. The isolated yield of monoester was 38% for melezitose and 54% for 1-kestose. The acylation of melezitose went preferentially to the $6^{\prime\prime}$ -OH of the glucose linked $1\rightarrow3$ to the fructose (69%) selectivity), to some extent to the $6'$ -OH of the fructose unit (22% selectivity) and a small proportion to the 6-OH of the other glucose residue (9% selectivity). In the case of 1-kestose, the esterification goes to the $6''$ -OH of the terminal fructose with high selectivity (90%), with only 7% to the $6'$ -OH of the middle fructose and 3% of other regioisomers. It is remarkable the strong preference for the fructofuranosyl ring compared with the results obtained with raffinose and melezitose.

The tetrasaccharide stachyose (4) was the last carbohydrate tested. We used slight modifications in the reaction conditions (see Table 1) in order to achieve complete solubility of this larger oligosaccharide. After 8 days the majority of the sugar remained unreacted (as seen by TLC) giving the lowest yield (26%) of isolated monoester amongst all the oligosaccharide tested. The regioselectivity partially correlates with that observed for raffinose; 79% of the esterification went to the 6-OH of the terminal galactose and 14% to the $6^{\prime\prime\prime}$ -OH of the fructose unit.

Transesterifications catalyzed by Thermomices lanuginosus lipase

Next, we studied the esterification of the selected oligosaccharides with the immobilized form of T. lanuginosus lipase on granulated silica (currently commercialized under the trade name Lipozyme TL IM). We selected this type of immobilization since it has higher selectivity to monoesters than other supported preparations when used in the synthesis of sucrose esters in two-solvent mixtures. 21 21 21 In addition, it has been reported^{[18](#page-115-0)} that lipases from T. lanuginosus and C. antarctica acylate sucrose in the same 2-methyl-2-butanol/DMSO mixtures. This observation lead us to use Lipozyme TL IM in the same tert-butanol/pyridine reaction mixtures described above for Novozym 435.

When we carried out the esterification reaction with Lipozyme TL IM for raffinose (1), the same regioisomer as for lipase Novozym 435 was obtained, but the isolated yield was much higher (79%) and the reaction time much shorter (24 h). Actually, the conversion of the reaction was 89% as determined by HPLC and no diester formation was detected (by TLC or HPLC) even at prolonged reaction times ([Fig. 1](#page-110-0)).

In the case of trisaccharides melezitose (2) and 1-kestose (3), similar reactivity was found for both sugars, although this was lower than for raffinose, requiring longer reaction time to reach the plateau of conversion ([Fig. 1\)](#page-110-0). The isolated yield of monoester after 72 h was 54% for melezitose and 57% for 1-kestose. Some starting material remained unreacted and only traces of diester were observed. The preferred position of acylation in the case of melezitose was the 6-OH of the glucose linked $1 \rightarrow 2$ to the fructose (61%) selectivity) while partial acylation occured at the primary hydroxyl of the other glucose unit (33% selectivity) and at the fructosyl 6'-OH (4% selectivity). On the other hand, 1-kestose was acylated at the 6-OH of the terminal glucose with 87% selectivity along with 13% of other unidentified regioisomers. This can be considered a high regioselectivity taking into account that 1-kestose has four primary hydroxyls (one more than raffinose) in its structure.

The esterification of the tetrasaccharide stachyose (4) with Lipozyme TL IM was also carried out using the same reaction conditions as with Novozym 435. Surprisingly, the initial reaction rate was very similar to the rate observed for 2 and 3 but after 3 days higher isolated yield of monoester was obtained (68%) with an excellent selectivity (96%)

Table 1. Acylation of tri- and tetrasaccharides using Novozym 435 and vinyl laurate as acylating agent

Carbohydrate	Solvent	Temperature $({}^{\circ}C)$	Reaction time(h)	Yield $(\%)$	Position of acylation	Selectivity $(\%)$
Raffinose	t -BuOH-pyridine (55/45)	60	72	48	Galactosyl 6-OH	> 99
Melezitose	t -BuOH-pyridine (55/45)	60	72	38	Glucosyl linked $1 \rightarrow 3$, 6 ⁿ -OH	69
					Fructosyl $6'$ -OH	22
					Glucosyl linked $1 \rightarrow 2$, 6-OH	
Kestose	t -BuOH-pyridine (55/45)	60	72	54	Terminal Fructosyl 6"-OH	90
					Middle Frutosyl 6'-OH	
Stachyose	t -BuOH-pyridine (50/50)	65	192	26	Galactosyl 6-OH	79
					Fructosyl $6^{\prime\prime\prime}$ -OH	14

Yields are referred to isolated monoester and selectivity corresponds to percentage of each regioisomer of saccharide monoester as found by HPLC/MS.

Figure 1. Kinetics of transesterification of the non-reducing oligosaccharides with vinyl laurate catalysed by Lipozyme TL IM. Conditions: (A) 0.04 M trisaccharide, 0.2 M vinyl laurate, 10 mg/ml biocatalyst, 10 mg/ml 3 Å molecular sieves, t-BuOH/pyridine (55:45), 60 °C. [raffinose (\circ), melezitose (\Box), 1-kestose (\Box)]. (B) 0.026 M tetrasaccharide, 0.131 M vinyl laurate, 8 mg/ml biocatalyst, 8 mg/ml 3 \AA molecular sieves, t-BuOH– pyridine (50/50), 65 °C [stachyose (\triangle)]. (C) 0.1 M trisaccharide, 0.5 M vinyl laurate 25 mg/ml biocatalyst, 25 mg/ml 3 Å molecular sieves, pyridine, 60° C [raffinose (\bullet)].

towards the 6-OH of the terminal galactose. A negligible amount of diacylated product was formed in this case.

T. lanuginosus lipase (immobilized on Celite) has previously shown better conversion than Novozym 435 in the acylation of sucrose with vinyl laurate, 18 what clearly agrees with the results we have obtained for the tri- and tetrasaccharides studied. It is also important to note that our isolated yields (54–79%) are higher than the previously reported in the literature^{[14](#page-115-0)} for the transesterification of maltotriose with vinyl laurate employing T. lanuginosus lipase (immobilized on Celite) in 2-methyl-2-butanol containing 20% DMSO, although we use longer reaction times and lower substrate concentration, what may be important for scale-up reactions. In any case, the tertbutanol/pyridine mixtures seem to be a good reaction media. In fact, we have observed that Lipozyme TL IM has significant activity in the transesterification of raffinose in pyridine (Fig. 1).

The regioselectivity we have found [\(Tables 1 and 2\)](#page-109-0) is different for both lipases (with the only exception of raffinose). Although the overall fold and catalytic machinery of the two enzymes are very similar, there are important structural differences that may explain the different behaviour. C antarctica lipase has a large binding pocket and likely no amphiphilic lid over it while in T. lanuginosus

lipase the binding pocket is less spacious and occluded by an helical lid.^{[22,23](#page-115-0)} Furthermore, computational studies of sucrose transesterification with vinyl laurate catalyzed by these lipases 24 have shown that not only the different binding pocket shape but also the different motion of the pocket during catalysis may explain the different regioselectivity observed with each lipase.

Transesterifications catalyzed by subtilisin Carlsberg

It has been reported that co-lyophilization of subtilisin Carlsberg with methyl- β -cyclodextrin (M β CD) improves the enzyme activity and enantioselectivity in organic solvents.^{[25](#page-115-0)} This form of the biocatalyst is superior to both the simple lyophilised powder and the cross-linked enzyme crystals (CLEC) preparations.^{[26](#page-115-0)} We decided to employ the M_bCD-subtilisin Carlsberg preparation in our oligosaccharide acylations expecting to obtain good yields with high levels of regioselectivity.

Subtilisin catalyzes carbohydrate acylation in DMF and pyridine; 12 12 12 although the former is a better solvent for oligosaccharides, the use of the amorphous form of the sugars allowed us to employ pyridine at 40° C facilitating the processing of the reaction. Using 0.2 M of raffinose (1) and 0.6 M of vinyl laurate in the presence of 22 mg/ml biocatalyst under vigorous stirring, high conversion was observed by TLC after 24 h and only traces of diester were noticed. Raffinose monoester was isolated in 74% yield with excellent regioselectivity (98%) towards the $1^{\prime\prime}$ -OH of the fructose moiety ([Table 3\)](#page-111-0).

Similar reaction conditions were employed for the remaining oligosaccharides. After 48 h, reactions were stopped when traces of diester were observed, although a small amount of starting material remained unreacted. Reaction over the tetrasaccharide stachyose (4) showed a great correlation with that of raffinose producing acylation in the same hydroxyl of the fructose unit (98% selectivity) and similar isolated yield (76%). On the other hand, acylation of melezitose and kestose went preferentially towards the $6'$ -OH of the central fructose in both cases.

The regioselectivity observed for stachyose is similar to the reported by Riva et al. 13 13 13 for acylation with trifluoroethyl butyrate in DMF, but for raffinose and melezitose our esterification conditions with vinyl laurate produce higher regioselectivity. Futhermore, we have obtained higher isolated yields by using $M\beta$ CD-subtilisin Carlsberg preparation than those described for carbohydrate acylation using a CLEC preparation of subtilisin^{[27](#page-115-0)} or a lyophilized

Table 2. Acylation of tri- and tetrasaccharides using Lipozyme TL IM and vinyl laurate as acylating agent

Carbohydrate	Solvent	Temperature $^{\circ}\textrm{C}$	Reaction time(h)	Yield $(\%)$	Position of acylation	Selectivity $(\%)$
Raffinose	t -BuOH-pyridine (55/45)	60	24	79	Galactosyl 6-OH	> 99
Melezitose	t -BuOH-pyridine (55/45)	60	72	54	Glucosyl linked $1 \rightarrow 2$, 6-OH	61
					Glucosyl linked $1 \rightarrow 3$, 6 ⁿ -OH	33
					Fructosyl 6'-OH	
Kestose	t -BuOH-pyridine (55/45)	60	72	57	Glucosyl 6-OH	87
Stachyose	t -BuOH-pyridine (50/50)	65	72	68	Galactosyl 6-OH	96
					Fructosyl $1^{\prime\prime\prime}$ -OH	

Yields are referred to isolated monoester and selectivity corresponds to percentage of each regioisomer of saccharide monoester as found by HPLC/MS.

Carbohydrate	Solvent	Temperature (°C)	Reaction time(h)	Yield $(\%)$	Position of acylation	Selectivity $(\%)$
Raffinose	Pyridine	40	24	74	Fructosyl 1"-OH	98
Melezitose	Pyridine	40	48	69	Fructosyl $6'$ -OH	73
					Glucosyl linked $1 \rightarrow 2$, 6-OH	16
					Glucosyl linked $1 \rightarrow 3$, 6 ⁿ -OH	6
Kestose	Pyridine	40	48	55	Middle Fructosyl 6'-OH	69
					Glucosyl 6-OH	11
					Terminal Fructosyl 6"-OH	10
Stachyose	Pyridine	40	48	76	Fructosyl 1 ^{/1} -OH	98

Table 3. Acylation of tri- and tetrasaccharides using M β CD-Subtilisin Carlsberg preparation and vinyl laurate as acylating agent

Yields are referred to isolated monoester and selectivity corresponds to percentage of each regioisomer of saccharide monoester as found by HPLC/MS.

subtilisin powder. 13 13 13 Computational studies of Subtilisin-catalyzed transesterification of sucrose^{[28](#page-115-0)} have shown that entropic factors are crucial in the enzyme regioselectivity. Modelling of the possible tetrahedral transition-state adducts find that in the 1[']OH-adduct the sucrose moiety is the most solvent exposed. This exposure confers a higher mobility upon the sucrose, which translates into a larger entropic stabilization. This approach also explains the results obtained with raffinose and stachyose as these compounds can be considered as sucrose derivatives modified at their glucosyl C-6 hydroxyl. It is probable than the 6'-OH adduct in both melezitose and 1-kestose is also the most solvent exposed.

In summary, we have carried out one-step acylation of triand tetrasaccharides catalyzed by hydrolases in good isolated yields. Key aspects in the reaction conditions are the use of the amorphous form of the saccharides and the specific solvent (tert-butanol/pyridine mixtures for lipases and pyridine for the protease). Moreover, it is the first time that these non-reducing oligosaccharides are acylated using lipases and are transesterified with a long chain acyl donor using subtilisin. High selectivity has been obtained for acylation with all the enzymes employed. This allows the access to a particular regioisomer by selecting the appropriate hydrolase, showing a picture of complementary regioselective acylation of the primary hydroxyl groups. At the same time, a correlation in the enzymatic behaviour towards the pair raffinose–stachyose is observed for all used hydrolases, and to a less extent for the pair 1-kestosemelezitose. Studies of the biological and physical properties of these new non-ionic surfactants are in progress.

3. Experimental

3.1. General methods

Anhydrous pyridine and vinyl laurate were supplied by Fluka; anhydrous *tert*-butanol, molecular sieves $(3 \text{ Å}, 8-12)$ mesh) and methyl- β -cyclodextrin from Aldrich; raffinose and melezitose from Sigma and stachyose and 1-kestose from TCI Chemicals. All the carbohydrates were used in their amorphous form prepared by lyophilization of the corresponding aqueous solutions. Granulated lipase from T. lanuginosus (Lipozyme TL IM), immobilized lipase from C. antarctica B (Novozym 435) and subtilisin Carslberg (purified powder) were kindly donated by Novozymes A/S. Molecular sieves were preactivated at ca. $350 \degree C$ for 12 h. All reactions were monitored by TLC on precoated Silica-Gel 60 plates (Alugram Sil G/UV $_{254}$ supplied by Macherey-Nagel), and detected by heating with Mostain (500 ml of 10% H₂SO₄, 25 g of $(NH_4)_6M_0T_7O_{24}$ 4H₂O, 1 g Ce(SO₄)₂ 4H₂O). The elution system was $CHCl₃$ –MeOH (2.5/1) for the reactions involving trisaccharides and EtOAc–MeOH–H₂O $(7/5/1)$ for the reactions with stachyose. Products were purified by flash chromatography with Aldrich Silica gel 60 (200–400 mesh) using a gradient of chloroform/methanol 5:1 to 2:1 (v/v) for the trisaccharides monolaurates and 5:1 to 1:1 v/v for the stachyose monolaurates.

NMR spectra were recorded on either a Bruker AVANCE 300 or ARX 400 [300 or 400 MHz (¹H) and 75 or 100 (¹³C)] at room temperature for solutions in $CD₃OD$. Chemical shifts are referred to the methanol multiplet, centered at 3.31 ppm for 1 H NMR and 49.0 ppm for 13 C NMR. Optical rotations (Sodium D line) were measured at 20° C with a Perkin-Elmer 241 for pure regioisomers (those oligosaccharide monolaurates obtained with $> 95\%$ regioselectivity). High resolution FAB $(+)$ mass spectral analyses were obtained on a Micromass AutoSpec-Q spectrometer. Infrared spectra were recorded using a Nicolet 20SXB FTIR spectrophotometer.

Analysis of the regioisomeric distribution of the isolated monoesters and the kinetic study of reactions catalyzed by Lipozyme TL IM were carried out by HPLC/MS. A Waters Alliance 2695 separation module was employed with a Waters Spherisorb 3 μ m ODS2 column (4.6 \times 250 mm) and a Waters Micromass ZQ mass spectrometer detector. The temperature of the column was set to 40 $^{\circ}$ C. Flow rate was 1.1 ml/min with splitting before the detection module. Mobile phases were acetonitrile/water mixtures in isocratic conditions. The ratio changed depending on the application as follows: for the analysis of the regioisomeric distribution of trisaccharide monolaurates, acetonitrile/water 35:65 (v:v) and for the more polar stachyose monolaurates, 30:70 (v:v); for the analysis of the kinetics of trisaccharide reactions, acetonitrile/water 45:55 (v:v) and for the stachyose reaction, 40:60 (v:v). Detection was done with positive ESI ionization in both Scan and SIR modes. For the kinetics studies, alliquots were concentrated to dryness, diluted with water and extracted with hexane to remove excess of vinyl laurate, prior to analysis.

3.2. General method for the transesterifications catalyzed by Novozym 435

Raffinose, melezitose or 1-kestose (202 mg, 0.4 mmol) were dissolved in anhydrous pyridine (4.5 ml) at 60° C before

careful addition of preheated (60 $^{\circ}$ C) anhydrous *tert*-butanol (5.5 ml) . Novozym 435 (100 mg) and 3 A molecular sieves (100 mg) were then added and the suspension maintained 30 min at 60° C with orbital shaking (250 rpm). Finally, vinyl laurate (457 mg, 2 mmol) was added. When conversion stopped by TLC, the mixture was cooled and filtered. The solvent was evaporated under vacuum at 45° C eliminating last traces of pyridine by co-evaporation with toluene. The remaining residue was subjected to flash chromatography. Concentration of pure fractions in vacuo afforded the monolaurates as amorphous white solids.

The tetrasaccharide stachyose (210 mg, 0.315 mmol) was dissolved in anhydrous pyridine (6 ml) at 65° C before careful addition of preheated (65 °C) anhydrous tert-butanol (6 ml). Novozym 435 (100 mg) and 3 Å molecular sieves (100 mg) were then added and the suspension maintained 30 min at 65° C with orbital shaking (250 rpm). Finally, vinyl laurate (360 mg) was added. The reaction was monitored and processed as described for trisaccharides.

3.2.1. 6-O-Lauroylraffinose (1a). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (132 mg, 48%). Only regioisomer 1a was obtained (HPLC/MS). $R_f = 0.34$; $[\alpha]_D$ $+80.7$ (c 7 in methanol); v_{max} (cm⁻¹) (KBr disks): 3410 br $(O-H)$, 1730 $(C=O)$; HRMS (FAB) : calcd for $C_{30}H_{54}O_{17}Na$ (M + Na⁺) 709.325871, found 709.324561; H NMR (CD₃OD, 400 MHz): δ 5.40 (d, 1H, $J_{1'-2'} = 3.7$ Hz, H-1'), 4.87 (d partially overlapped with residual water, 1H, H-1), 4.24 (m, 1H, H-6a), 4.22 (dd, 1H, $J_{6a-b} = 11.3$ Hz, J_{6b-5} = 2.6 Hz, H-6b), ca. 4.10 (m, 2H, H-3ⁿ, H-4ⁿ); ca. 4.04 $(m, 2H, H-5, H-5')$, 3.88 (br.d, 1H, $J=3.0$ Hz, H-4), 3.83 (dd, 1H, $J_{6a-b} = 11.1$ Hz, $J_{6a-5'} = 6.1$ Hz, H-6^ta), ca. 3.81 $(m, 1H, H-3)$, ca. 3.76 $(m, 3H, H-5)$, H6ⁿa, H6ⁿb), ca. 3.73 (m, 1H, H-2), 3.71 (t, 1H, $J_{2'-3'} = J_{3'-4'} = 9.4$ Hz, H-3[']) ca. 3.70 (m, 1H, H-6^tb), 3.64 (d, 1H, $J_{1''a-b} = 12.3$ Hz, H-1^ta), 3.60 (d, 1H, $J_{1^n a-b} = 12.3$ Hz, H-1ⁿb), 3.43 (dd, 1H, $J_{2^{\prime}-3^{\prime}} =$ 9.7 Hz, $J_{1'-2'}=3.7$ Hz, H-2[']), 3.26 (t, 1H, $J_{3'-4'}=J_{4'-5'}=$ 9.5 Hz, H-4⁷), 2.34 (t, 2H, J=7.4 Hz, -CH₂-CO-), 1.61 (m, 2H, CH₂–CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃-); ¹³C NMR (CD₃OD, 100 MHz): δ 175.4 (C=O), 105.3 (C-2"), 100.5 (C-1), 93.4 (C-1'), 83.5 $(C-5'')$, 79.1 $(C-3'')$, 75.4 $(C-4'')$, 74.4 $(C-3')$, 73.2 $(C-5')$, 73.1 (C-2'), 72.1 (C-4'), 71.1 (C-3), 70.9 (C-4), 70.3 (C-2), 69.9 (C-5), 68.5 (C-6'), 64.9 (C-6), 64.2 (C-1"), 63.3 (C-6"), 35.0 (–CH2–CO–), 33.1, 30.7, 30.6, 30.5, 30.4, 30.2, 26.0, 23.7 ($-CH_2$ -lauroyl backbone), 14.5 (CH_3 -lauroyl).

3.2.2. $6''$ -O-Lauroylmelezitose (2a). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (104 mg, 38%). R_f = 0.63; v_{max} (cm⁻¹) (KBr disks): 3400 br (O-H), 1725 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ (M+Na⁺) 709.325871, found 709.326346. Regioisomeric proportion $6''/6'/6 = 69/22/9$ (HPLC/MS). NMR assignments of main regioisomer 2a ¹H NMR (CD₃OD, 400 MHz): δ 5.47 (d, 1H, J_{1-2} = 3.8 Hz, H-1), 5.08 (d, 1H, $J_{1''-2''}$ = 3.6 Hz, H-1ⁿ), 4.42 (dd, 1H, $J_{6''a-b} = 11.6$ Hz, $J_{6''a-5''} = 1.5$ Hz, H-6ⁿa), 4.29 (t, 1H, $J_{3'-4'} = J_{4'-5'} = 7.7$ Hz, H-4'), 4.24 (dd, 1H, $J_{6''a-b} =$ 11.6 Hz, $J_{6''b-5''}=6.4$ Hz, H_{-6} ⁿb), 4.19 (d, 1H, $J_{3'-4'}=$ 7.7 Hz, H-3[']), 4.11 (m, 1H, H-5^{''}), 3.91 (m, 1H, H-5), 3.85 (dd, 1H, J_{6a-b} =11.9 Hz, J_{6a-5} =1.4 Hz, H-6a), ca. 3.78 (m,

2H, H6'a, H6'b), ca. 3.73 (m, 1H, H-5') ca. 3.72 (m, 1H, H1[']a), 3.70 (dd, 1H, $J_{6a-b} = 11.9$ Hz, $J_{6b-5} = 4.9$ Hz, H-6b), 3.69 (t, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=J_{3^{\prime\prime}-4^{\prime\prime}}=9.6$ Hz, H-3^{$\prime\prime$}), 3.62 (t, 1H, $J_{2-3}=J_{3-4}=9.6$ Hz, H-3), 3.58 (d, 1H, H-1^tb), 3.43 (dd, 1H, $J_{2-3}=9.7$ Hz, $J_{1-2}=3.8$ Hz, H-2), 3.41 (dd, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=$ 9.7 Hz, $J_{1^{\prime\prime}-2^{\prime\prime}}$ = 3.8 Hz, H-2^{$\prime\prime$}), 3.35 (t, 1H, $J_{3^{\prime\prime}-4^{\prime\prime}}$ = $J_{4^{\prime\prime}-5^{\prime\prime}}$ = 9.6 Hz, H-4^{$\prime\prime$}), 3.31 (t, 1H, $J_{3-4}=J_{4-5}=9.6$ Hz, H-4), 2.37 (t, 2H, $J=7.5$ Hz, $-CH_2-CO-$), 1.61 (m, 2H, CH_2-CH_2- CO–), 1.29 (m, 16H, $-CH_2$ – chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃–); ¹³C NMR (CD₃OD, 100 MHz): δ 175.5 (C=O), 105.3 (C-2'), 101.6 (C-1"), 93.2 (C-1), 85.7 (C-3'), 83.7 (C- $5'$), 75.1 (C-3), 74.8 (C-3ⁿ, C-4¹), 74.2 (C-5), 73.6 (C-2ⁿ), 73.2 (C-2), 71.8 (C-4^{$\prime\prime$}), 71.7 (C-5^{$\prime\prime$}, C-4), 64.8 (C-6^{$\prime\prime$}), 64.6 $(C-1)$, 63.3 $(C-6)$, 62.6 $(C-6)$, 35.0 $(-CH_2-CO-)$, 33.1, 30.7, 30.6, 30.4, 30.2, 26.0, 23.7 (-CH₂-lauroyl backbone), 14.4 ($CH₃$ -lauroyl).

3.2.3. $6''$ -O-Lauroyl(1-kestose) (3a). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (148 mg, 54%). $R_f = 0.33$; v_{max} (cm⁻¹) (KBr disks): 3400 br (O–H), 1730 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ (M+ $Na⁺$) 709.325871, found 709.324767. Main regioisomer 3a was produced with 90% regioselectivity (HPLC/MS). NMR assignments of $3a⁻¹H NMR (CD₃OD, 400 MHz): \delta 5.38$ (d, 1H, J_{1-2} = 3.8 Hz, H-1), 4.41 (dd, 1H, $J_{6''a-b}$ = 11.8 Hz, $J_{6''a-5''}=8.0$ Hz, H-6ⁿa), 4.23 (dd, 1H, $J_{6''a-b}=11.8$ Hz, $J_{6''b-5''}=3.0$ Hz, H-6ⁿb), 4.19 (d, 1H, $J_{3'-4'}=8.4$ Hz, H-3^t), 4.12 (d, 1H, $J_{3''-4''}=8.1$ Hz, H-3ⁿ), 4.02 (m, 1H, H4¹), 4.00 (m, 1H, H4^{j'}), 3.91 (td, 1H, $J_{4''-5''}=J_{5''-6''a}=7.9$ Hz, $J_{5''-6''b}$ = 3.0 Hz, H-5^{''}), 3.81 (m, 2H, H-5, H1[']a), ca. 3.77 (m, 1H, H-1'b), ca. 3.76 (m, 3H, H-5', H6'a, H-6'b), ca. 3.71 (m, 2H, H-6a, H-6b), 3.67 (t, 1H, $J_{2-3}=J_{3-4}=9.4$ Hz, H-3), 3.65 (d, 1H, $J_{1''a-b}$ =12.0 Hz, H-1ⁿa), 3.58 (d, 1H, $J_{1''a-b}$ = 12.0 Hz, H-1ⁿb), 3.38 (dd, 1H, J_{2-3} =9.6 Hz, J_{1-2} =3.8 Hz, H-2), 3.34 (t, 1H, $J_{3-4} = J_{4-5} = 9.4$ Hz, H-4), 2.36 (t, 2H, $J =$ 7.5 Hz, $-CH_2$ –CO–), 1.61 (m, 2H, CH_2 –CH₂–CO–), 1.29 (m, 16H, $-CH_2$ – chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃–); ¹³C NMR (CD₃OD, 100 MHz): δ 175.5 (C=O), 105.6 $(C-2'')$, 104.9 $(C-2')$, 94.1 $(C-1)$, 83.6 $(C-5')$, 80.7 $(C-5'')$, 79.5 (C-3'), 78.6 (C-3"), 77.3 (C-4"), 75.6 (C-4'), 74.8 $(C-3)$, 74.4 $(C-5)$, 73.3 $(C-2)$, 71.5 $(C-4)$, 66.8 $(C-6'')$, 63.2 (C-6'), 62.9 (C-1'), 62.3 (C-6), 61.9 (C-1"), 35.0 $(-CH₂-CO₋), 33.1, 30.7, 30.6, 30.4, 30.2, 26.0, 23.7, (-CH₂-C)$ lauroyl backbone), 14.4 ($CH₃$ -lauroyl).

3.2.4. 6-O-Lauroylstachyose (4a). The general procedure outlined above was followed. After 8 days the reaction was stopped and the monoester isolated (69 mg, 26%). Regioisomeric proportion $6/6''' = 79/14\% + 7\%$ other regioisomers (HPLC/MS). See characterization of main regioisomer 4a in next section.

3.3. General method for the transesterifications catalyzed by Lipozyme TL IM

Exactly the same method as the Novozym 435 catalyzed reactions was used changing just the biocatalyst to Lipozyme TL IM.

3.3.1. 6-O-Lauroylraffinose (1a). The general procedure outlined above was followed. After 24 h the reaction was stopped and the monoester isolated (217 mg, 79%).

Only one regioisomer 1a was obtained (HPLC/MS), which is the same obtained with Novozym 435 catalysis.

3.3.2. 6-O-Lauroylmelezitose (2b). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (148 mg, 54%). R_f = 0.33; v_{max} (cm⁻¹) (KBr disks): 3400 br (O-H), 1725 (C=O); HRMS (FAB): calcd for C₃₀H₅₄O₁₇Na (M+Na⁺) 709.325871, found 709.326346. Regioisomeric proportion $6/6''/6' = 61/33/4\% + 2\%$ other regioisomers (HPLC/MS). NMR assignments of main regioisomer $2\mathbf{b}^{-1}H$ NMR (CD₃OD, 400 MHz): δ 5.42 (d, 1H, $J_{1-2}=3.8$ Hz, H-1), 5.08 (d, 1H, $J_{1''-2''}=3.8$ Hz, H-1^{''}), 4.44 (dd, 1H, $J_{6a-b}=$ 12.0 Hz, $J_{6a-5} = 1.4$ Hz, H-6a), 4.16 (dd, 1H, $J_{6a-b} =$ 12.0 Hz, $J_{6b-5} = 5.6$ Hz, H-6b), ca. 4.23 (m, 1H, H-4⁷), ca. 4.20 (m, 1H, H-3'), 4.04 (m, 1H, H-5), 3.97 (m, 1H, H-5"), ca. 3.91 (m, 1H, $H-6''a$), ca. 3.87 (m, 1H, $H-1'a$), ca. 3.85 (m, 1H, H-5[']), ca. 3.70 (m, 2H, H-6[']a, H-6[']b), 3.69 (t, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=J_{3^{\prime\prime}-4^{\prime\prime}}=9.3$ Hz, H-3^{$\prime\prime$}), 3.65 (dd, 1H, $J_{6^{\prime\prime}a-b}=$ 12.0 Hz, $J_{6''b-5''}=5.2$ Hz, H-6ⁿb), 3.61 (t, 1H, $J_{2-3}=$ $J_{3-4} = 9.3 \text{ Hz}, \text{ H-3}, 3.57 \text{ (d, 1H, } J_{1\text{a}-\text{b}} = 12.1 \text{ Hz}, \text{ H-1}^{\prime} \text{b}),$ 3.43 (dd, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=9.8$ Hz, $J_{2^{\prime\prime}-1^{\prime\prime}}=3.7$ Hz, H-2^{$\prime\prime$}), 3.41 (dd, 1H, $J_{2-3}=9.8$ Hz, $J_{2-1}=3.9$ Hz, H-2), 3.27 (t, 1H, $J_{3''-4''}=J_{4''-5''}=9.4$ Hz, H-4ⁿ), 3.26 (t, 1H, $J_{3-4}=J_{4-5}=$ 9.4 Hz, H-4), 2.37 (t, 2H, $J=7.5$ Hz, $-CH_2$ –CO–), 1.62 (m, 2H, CH₂–CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃-); ¹³C NMR (CD₃OD, 100 MHz): δ 175.5 (C=O), 105.4 (C-2'), 102.2 (C-1^{π}), 93.2 (C-1), 86.1 $(C-3')$, 83.4 $(C-5')$ 75.0 $(C-3)$, 74.7 $(C-3'')$, 74.6 $(C-4')$, 73.9 $(C-5'')$, 73.7 $(C-2'')$, 73.1 $(C-2)$, 72.1 $(C-4'')$, 71.9 $(C-5)$, 71.6 $(C-4)$, 64.8 $(C-6)$, 64.0 $(C-1', C-6')$, 62.9 $(C-6'')$, 34.9 (–CH2–CO–), 33.1, 30.8, 30.7, 30.5, 30.4, 30.3, 26.0, 23.7 (–CH2-lauroyl backbone), 14.5 (CH3-lauroyl).

3.3.3. 6-O-Lauroyl(1-kestose) (3b). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (157 mg, 57%). R_f = 0.33; v_{max} (cm⁻¹) (KBr disks): 3400 br (O–H), 1730 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ (M+Na⁺) 709.325871, found 709.324767. Main regioisomer 3b was produced with 87% regioselectivity (HPLC/MS). NMR assignments for main regioisomer $3b$ ¹H NMR (CD₃OD, 400 MHz): δ 5.39 (d, 1H, J_{1-2} = 3.8 Hz, H-1), 4.39 (dd, 1H, $J_{6a-b} = 12.0$ Hz, $J_{6a-5} = 1.5$ Hz, H-6a), 4.20 (d, 1H, $J_{3/4} =$ 8.4 Hz, H-3'), 4.18 (dd, 1H, $J_{6a-b} = 12.0$ Hz, $J_{6b-5} = 5.0$ Hz, H-6b), 4.13 (d, 1H, $J_{3''-4''}=8.3$ Hz, H-3ⁿ), ca. 4.01 (m, 1H, $H-4''$), 4.00 (m, 1H, H-5), ca. 3.99 (m, 1H, H-4'), 3.82 (d, $1H, J_{1a-b} = 10.4 \text{ Hz}, H-1a$, ca. 3.77 (m, 1H, H1^tb), ca. 3.76 (m, 1H, H5'), ca. 3.75 (m, 4H, H6'a, H6'b, H6"a, H6"b), ca. 3.74 (m, 1H, H-5"), 3.69 (t, 1H, $J_{2-3}=J_{3-4}=9.4$ Hz, H-3), 3.64 (d, 1H, $J_{1^n a-b}$ =12.0 Hz, H-1ⁿa), 3.59 (d, 1H, $J_{1^n a-b}$ = 12.0 Hz, H-1["]b), 3.39 (dd, 1H, J_{2-3} =9.7 Hz, J_{1-2} =3.8 Hz, H-2), 3.30 (t, 1H, $J_{3-4} = J_{4-5} = 9.4$ Hz, H-4), 2.37 (t, 2H, $J =$ 7.5 Hz, $-CH_2$ -CO–), 1.61 (m, 2H, CH_2 -CH₂-CO–), 1.29 (m, 16H, -CH₂- chain), 0.90 (t, 3H, $J=6.7$ Hz, CH₃-); ¹³C NMR (CD₃OD, 100 MHz): δ 175.5 (C=O), 105.3 (C-2[']), 104.9 (C-2^{$\prime\prime$}), 93.8 (C-1), 83.7 (C-5^{\prime}), 83.4 (C-5^{$\prime\prime$}), 79.2 (C-3'), 78.7 (C-3"), 76.3 (C-4"), 75.7 (C-4'), 74.5 (C-3), 73.2 $(C-2)$, 72.0 $(C-5)$, 71.6 $(C-4)$, 64.6 $(C-6)$, 63.9 $(C-6)$, 63.7 $(C-6^{\prime})$, 63.0 $(C-1^{\prime})$, 62.3 $(C-1^{\prime\prime})$, 34.9 $(-CH_2-CO-)$, 33.1, 30.7, 30.6, 30.5, 30.2, 26.0, 23.7 (–CH₂-lauroyl backbone), 14.4 ($CH₃$ -lauroyl).

3.3.4. 6-O-Lauroylstachyose (4a). The general procedure outlined above was followed. After 3 days the reaction was stopped and the monoester isolated (183 mg, 68%). Regioisomer 4a was obtained with $> 95\%$ regioselectivity (HPLC/MS). $R_f = 0.63$; $[\alpha]_D + 106.9$ (c 7 in methanol); v_{max} $(cm⁻¹)$ (KBr disks): 3410 br (O–H), 1730 (C=O); HRMS (FAB): calcd for $C_{36}H_{64}O_{22}Na$ (M+Na⁺) 871.378694, found 871.378528; ¹H NMR (CD₃OD, 400 MHz): δ 5.41 (d, 1H, $J_1''_{-2} = 3.7$ Hz, H-1ⁿ), 4.86 (d, 1H, H-1), 4.85 (d, 1H, $J_{1'-2'} = 3.6$ Hz, H-1'), 4.27 (dd, 1H, $J_{6a-b} = 11.2$ Hz, $J_{6a-5} =$ 4.7 Hz, H-6a), 4.21 (dd, 1H, $J_{6a-b} = 11.2$ Hz, $J_{6b-5} = 7.5$ Hz, H-6b), 4.12 (m, 1H, H-5), ca. 4.10 (m, 2H, H-3 $''$, H-4 $''$), ca. 4.05 (m, 1H, H-5"), ca. 4.04 (m, 1H, H-5'), 3.90 (m, 1H, H-4), 3.88 (dd, 1H, 1H, $J_{6''a-b} = 11.0$ Hz, $J_{6''a-5''} = 5.9$ Hz, $H-6''a$), ca. 3.87 (m, 1H, $H-4'$), 3.85 (m, 1H, $H-6'a$), 3.83 (m, 1H, H-3), 3.81 (m, 1H, H-3'), ca. 3.77 (m, 3H, H-6^ma, $H-6'''$ b, H-2), ca. 3.76 (m, 1H, H-5^{$'''$}), ca. 3.75 (m, 1H, H-2[']), 3.71 (m, 2H, H-3ⁿ, H-6ⁿb), 3.63 (d, 1H, $J_1\text{m}_{a-b}=12.4$ Hz, H-1^{*m*}a), 3.62 (m, 1H, H-6^{*l*}b), 3.60 (d, 1H, J_1 _{*m*}_{a-b} = 12.4 Hz, H-1^mb), 3.44 (dd, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=9.7$ Hz, $J_{1^{\prime\prime}-2^{\prime\prime}}=3.7$ Hz, H-2ⁿ), 3.29 ((t, 1H, $J_{3^{\prime\prime}-4^{\prime\prime}}=J_{4^{\prime\prime}-5^{\prime\prime}}=9.5$ Hz, H-4ⁿ), 2.37 (t, $2H, J=7.5$ Hz, $-CH_2$ –CO–), 1.62 (m, 2H, CH₂–CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃–); ¹³C NMR (CD₃OD, 100 MHz): δ 175.4 (C=O), 105.3 $(C-2^{\prime\prime\prime})$, 100.4 $(C-1^{\prime})$, 99.8 $(C-1)$, 93.4 $(C-1^{\prime\prime})$, 83.5 $(C-5^{\prime\prime\prime})$, 79.1 $(C-3^{\prime\prime\prime})$, 75.3 $(C-4^{\prime\prime\prime})$, 74.5 $(C-3^{\prime\prime})$, 73.1 $(C-5^{\prime\prime})$, 73.0 $(C-2'')$, 72.1 $(C-4'')$, 71.3 $(C-3')^a$, 71.2 $(C-3)^a$, 71.0 $(C-4)$, $C-4'$), 70.6 $(C-2)^b$, 70.5 $(C-5')^b$, 70.1 $(C-2')^b$, 69.9 $(C-5)$, 68.4 (C-6"), 68.0 (C-6'), 64.9 (C-6), 64.2 (C-1"'), 63.2 $(C-6^{\prime\prime\prime})$, 35.0 ($-CH_2$ –CO–), 33.1, 30.8, 30.7, 30.5, 30.3, 26.0, 23.7 (-CH₂-lauroyl backbone), 14.5 (CH₃-lauroyl). Note: ¹³C NMR assignments marked with the same letter may be interchangeable.

3.4. General method for the transesterifications catalyzed by subtilisin Carslberg

 $M\beta$ CD-subtilisin Carlsberg preparation was obtained as previously described^{[26](#page-115-0)} using 0.1 M KOH instead of phosphate buffer for the pH adjustment in order to prevent side alkaline catalysis in the transesterificatin reactions, which would promote competitive non-regioselective acylations and even more substitution.^{[29](#page-115-0)}

In a typical experiment a solution of raffinose, melezitose or 1-kestose (125 mg, 0.25 mmol) in anhydrous pyridine (2 ml) and vinyl laurate (170 mg, 0.74 mmol) were shaken with vigorous magnetic stirring at 40° C in the presence of $M\beta$ CD-subtilisin Carlsberg (45 mg). When conversion stopped by TLC, the mixture was cooled and filtered. The solution was concentrated under vacuum at 45° C eliminating last traces of pyridine by co-evaporation with toluene. The remaining residue was subjected to flash chromatography. Concentration of pure fractions in vacuo afforded the monolaurates as amorphous white solids.

For the tetrasaccharide transesterification, the enzyme (70 mg), stachyose (150 mg, 0.225 mmol) and vinyl laurate (156 mg, 0.68 mmol) were mixed in anhydrous pyridine (2 ml). The same reaction conditions and work-up described for the trisaccharides was followed.

3.4.1. $1^{\prime\prime}$ -O-Lauroylraffinose (1b). The general procedure outlined above was followed. After 24 h the reaction was stopped and the monoester isolated (127 mg, 74%). Regioisomer 1b was obtained with 98% regioselectivity (HPLC/ MS). $R_f = 0.30$; $[\alpha]_D + 94$ (c 7 in methanol); ν_{max} (cm⁻¹) (KBr disks): 3410 br (O-H), 1735 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ $(M+Na^{+})$ 709.325871, found 709.326289; ¹H NMR (CD₃OD, 300 MHz): δ 5.41 (d, 1H, $J_{1'-2'}=3.8$ Hz, H-1'), 4.89 (d, 1H, $J_{1-2}=3.2$ Hz, H-1), 4.35 (d, 1H, $J_{1^n a-b}$ = 12.2 Hz, H-1ⁿa), 4.12 (d, 1H, $J_{1^n a-b}$ = 12.2 Hz, H-1 $\frac{1}{1}$ b), 4.10 (m, 1H, H-4 $\frac{1}{1}$), 4.09 (m, 1H, H-3 $\frac{1}{1}$), 4.05 (m, 1H, H-5[']), 3.91 (m, 1H, H-3), 3.87 (dd, 1H, $J_{6a-b} = 11.5$ Hz, J_{6a-5} = 6.0 Hz, H-6^ta), 3.85 (m, 1H, H-5), 3.78 (m, 1H, H-4), ca. 3.77 (m, 1H, H-2), ca. 3.75 (m, 2H, H-6"a, H-6"b), ca. 3.75 (m, 1H, H6'-b), ca. 3.73 (m, 1H, H-5"), 3.70 (m, 2H, H-6a, H-6b), 3.69 (t, 1H, $J_{2'-3'} = J_{3'-4'} = 9.4$ Hz, H-3[']), 3.43 (dd, 1H, $J_{2'-3'}=9.6$ Hz, $J_{1'-2'}=3.8$ Hz, H-2'), 3.32 (t, 1H, $J_{3'-4'}=$ $J_{4'-5'}=9.2$ Hz, $H-4^7$), 2.37 (t, 2H, $J=7.4$ Hz, $-CH_2$ -CO-), 1.62 (m, 2H, CH₂–CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, $J=6.7$ Hz, CH_3 –); ¹³C NMR (CD₃OD, 75 MHz): δ 174.8 (C=O), 104.4 (C-2"), 100.5 (C-1), 93.9 (C-1'), 83.6 $(C-5''), 78.6 (C-3''), 74.7 (C-4''), 74.4 (C-3'), 73.3 (C-5'), 72.8$ (C-2[']), 72.4 (C-5), 72.0, (C-4[']), 71.4 (C-3), 71.0 (C-4), 70.5 $(C-2)$, 68.3 $(C-6)$, 64.1 $(C-1'')$, 63.0 $(C-6'')$, 62.8 $(C-6)$, 35.0 $(-CH₂-CO₋)$, 33.0, 30.7, 30.6, 30.4, 30.2, 26.0, 23.7 $(-CH₂$ lauroyl backbone), 14.4 ($CH₃$ -lauroyl).

3.4.2. $6'$ - O -Lauroylmelezitose (2c). The general procedure outlined above was followed. After 48 h the reaction was stopped and the monoester isolated (119 mg, 69%). R_f = 0.32; ν_{max} (cm⁻¹) (KBr disks): 3400 br (O–H), 1725 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ (M+Na⁺) 709.325871, found 709.325243. Regioisomeric proportion $6′/6/6″ = 73/16/6% + 5%$ other regioisomers (HPLC/MS). NMR assignments of main regioisomer $2c$, ¹H NMR (CD₃OD, 300 MHz): δ 5.41 (d, 1H, $J_{1-2} = 3.8$ Hz, H-1), 5.12 (d, 1H, $J_{1''-2''}=3.7$ Hz, H-1ⁿ), 4.43 (dd, 1H, $J_{6'ab}=$ 11.8 Hz, $J_{6a-5} = 7.7$ Hz, H-6^ta), 4.35 (dd, 1H, $J_{6a-b} =$ 11.8 Hz, $J_{6a-5'} = 3.7$ Hz, H-6^tb), 4.34 (t, 1H, $J_{3'-4'} = J_{4'-5'} =$ 8.1 Hz, H-4^T), 4.23 (d, 1H, $J_{3'-4'}=8.1$ Hz, H-3^T), 3.95 (m, 1H, H-5[']), ca. 3.90 (m, 2H, H-5, H-5^{''}), ca. 3.88 (m, 2H, H-6a, H-6ⁿa), 3.82 (d, 1H, $J_{1a-b} = 12.3$ Hz, H-1¹a), ca. 3.69 (m, 1H, H-3^{$\prime\prime$}), ca. 3.68 (m, 2H, H-6b, H-6 $\prime\prime$ b), ca. 3.63 (m, 1H, H-3), 3.60 (d, 1H, $J_{1/a-b} = 12.3$ Hz, H-1[']b), 3.42 (dd, 1H, H-2^{*i*'}), 3.40 (dd, 1H, H-2), ca. 3.28 (m, 2H, H-4, H-4"), 2.35 (t, 2H, $J=$ 7.5 Hz, $-CH_2$ –CO–), 1.62 (m, 2H, CH_2 –CH₂–CO–), 1.29 (m, 16H, $-CH_2$ – chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃–); ¹³C NMR $(CD_3OD, 75 MHz)$: δ 175.5 $(C=O), 105.6$ $(C-2'), 101.7$ $(C-1¹)$, 93.1 $(C-1)$, 85.2 $(C-3¹)$, 80.4 $(C-5¹)$, 75.7 $(C-4¹)$, 75.1 $(C-3)$, 74.9 $(C-3'')$, 74.0 $(C-5, C-5')$, 73.7 $(C-2'')$, 73.3 $(C-2)$, 72.1 (C-4"), 72.0 (C-4), 66.6 (C-6'), 64.2 (C-1'), 62.9 (C-6"), 62.8 (C-6), 35.0 (–CH2–CO–), 33.1, 30.7, 30.6, 30.4, 30.2, 26.0, 23.7 ($-CH_2$ -lauroyl backbone), 14.4 (CH_3 -lauroyl).

3.4.3. 6'-O-Lauroyl(1-kestose) (3c). The general procedure outlined above was followed. After 48 h the reaction was stopped and the monoester isolated (94 mg, 55%). R_f = 0.33; v_{max} (cm⁻¹) (KBr disks): 3400 br (O–H), 1730 (C=O); HRMS (FAB): calcd for $C_{30}H_{54}O_{17}Na$ (M+Na⁺) 709.325871, found 709.324767. Main regioisomer 3c was produced with 69% regioselectivity (HPLC/MS). NMR assignments for main regioisomer $3c$: ¹H NMR (CD₃OD, 400 MHz): δ 5.37 (d, 1H, J_{1-2} = 3.8 Hz, H-1), 4.31 (m, 2H,

H6^{\prime}a, H6^{\prime}b), 4.20 (d, 1H, $J_{3/4}$ = 8.3 Hz, H-3^{\prime}), 4.13 (d, 1H, $J_{3''-4''} = 8.1$ Hz, H-3ⁿ), 4.01 (m, 1H, H-4ⁿ), 3.99 (m, 1H, H-4¹), 3.89 (m, 1H, H-5[']), 3.86 (d, 1H, $J_{1a-b} = 11.8$ Hz, H-1[']a), 3.80 (m, 1H, H-5), ca. 3.74 (m, $3H$, $H-5$ ⁿ, $H6$ ⁿa, $H6$ ⁿb), 3.70 (m, 3H, H-1'b, H-6a, H-6b), 3.67 (t, 1H, $J_{2-3}=J_{3-4}=$ 9.6 Hz, H-3), 3.65 (d, 1H, $J_{1''a-b}$ = 11.9 Hz, H-1ⁿa), 3.58 (d, 1H, $J_{1^n a-b} = 11.9$ Hz, H-1ⁿb), 3.38 (dd, 1H, $J_{2-3} = 9.7$ Hz, $J_{1-2}=3.8$ Hz, H-2), 3.33 (t, 1H, $J_{3-4}=J_{4-5}=9.6$ Hz, H-4), 2.37 (t, 2H, $J=7.5$ Hz, $-CH_2$ –CO–), 1.61 (m, 2H, CH₂– CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, J= 6.7 Hz, CH₃-); ¹³C NMR (CD₃OD, 100 MHz): δ 175.5 $(C=0)$, 105.3 $(C-2'')$, 105.2 $(C-2')$, 94.0 $(C-1)$, 83.5 $(C-5'')$, 80.5 (C-5'), 78.8 (C-3'), 78.7 (C-3"), 76.5 (C-4'), 76.3 (C-4"), 74.8 (C-3), 74.2 (C-5), 73.4 (C-2), 71.5 (C-4), 66.5 (C- $6'$), 63.9 (C-6^{*ii*}), 62.6 (C-1^{*i*}), 62.4 (C-1^{*ii*}), 62.3 (C-6), 34.9 $(-CH₂-CO₋), 33.1, 30.8, 30.6, 30.5, 30.2, 26.0, 23.7 (-CH₂-C)$ lauroyl backbone), 14.4 ($CH₃$ -lauroyl).

3.4.4. $1^{\prime\prime\prime}$ -*O*-Lauroylstachyose (4b). The general procedure outlined above was followed. After 48 h the reaction was stopped and the monoester isolated (145 mg, 76%). Regioisomer 4b was obtained with 98% regioselectivity (HPLC/MS). $R_f = 0.63$; $[\alpha]_D + 105.4$ (c 7 in methanol); ν_{max} $(cm⁻¹)$ (KBr disks): 3410 br (O–H), 1735 (C=O); HRMS (FAB): calcd for $C_{36}H_{64}O_{22}Na$ (M+Na⁺) 871.378694, found 871.378598; ¹H NMR (CD₃OD, 400 MHz): δ 5.41 (d, 1H, $J_{1''-2''}=3.7$ Hz, H-1^{$\prime\prime$}), 4.88 (d, 1H, $J_{1-2}=3.2$ Hz, H-1), 4.86 (d, 1H, $J_{1'-2'}=3.7$ Hz, H-1[']), 4.35 (d, 1H, $J_{1''a-b} =$ 12.1 Hz, H-1^{m}a), 4.11 (d, 1H, $J_1m_{a-b}=12.1$ Hz, H-1 m b), ca. 4.08 (m, 2H, H-3^{III}, H-4^{III}), 4.07 (m, 1H, H-5¹), 4.05 (m, 1H, H-5ⁿ), 3.93 (m, 1H, H-4^t), 3.91 (dd, 1H, $J_{6^n a-b} = 11.1$ Hz, $J_{6''a-5''}=5.6$ Hz, H-6^{n}a), ca. 3.89 (m, 1H, H-5), ca. 3.85 (m, 1H, H-6'a), 3.83 (m, 1H, H-3), 3.80 (m, 1H, H-4), ca. 3.76 $(m, 4H, H-2, H-2', H-6^{III}a, H-6^{III}b), ca. 3.73 (m, 1H, H-3'),$ ca. 3.72 (m, 1H, H-5^m), ca. 3.71 (m, 2H, H-6a, H-6b), 3.69 (m, 1H,, H-6ⁿb), 3.67 (t, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}}=J_{3^{\prime\prime}-4^{\prime\prime}}=9.5$ Hz, H-3ⁿ), ca. 3.65 (m, 1H, H-6^tb), 3.43 (dd, 1H, $J_{2^{\prime\prime}-3^{\prime\prime}} = 9.7 \text{ Hz}$, $J_{1^{\prime\prime}-2^{\prime\prime}} = 3.7$ Hz, H-2^{$\prime\prime$}), 3.32 (t, 1H, $J_{3^{\prime\prime}-4^{\prime\prime}} = \tilde{J}_{4^{\prime}-5^{\prime}} = 9.5$ Hz, H-4^{$\prime\prime$}), 2.37 (t, 2H, J=7.5 Hz, –CH₂–CO–), 1.63 (m, 2H, CH_2 –CH₂–CO–), 1.29 (m, 16H, –CH₂– chain), 0.90 (t, 3H, $J=6.8$ Hz, CH₃–); ¹³C NMR (CD₃OD, 60 MHz): δ 174.8 $(C=0)$, 104.1 $(C-2^{\prime\prime\prime})$, 100.4 $(C-1^{\prime})$, 100.1 $(C-1)$, 93.9 $(C-1'')$, 83.7 $(C-5''')$, 78.6 $(C-3''')$, 74.7 $(C-4''')$, 74.5 $(C-3'')$, 73.1 (C-5"), 72.8 (C-2"), 72.4 (C-5), 72.0 (C-4"), 71.5 $(C-3)^{a}$, 71.3 $(C-3')^{a}$, 71.0 $(C-4)^{b}$, 70.9 $(C-4')^{b}$, 70.5 $(C-5')^{c}$, 70.2 (C-2, C-2')^c, 68.2 (C-6"), 67.8 (C-6'), 64.0 (C-1^m), 63.1 $(C-6^{\prime\prime\prime})$, 62.6 $(C-6)$, 35.0 $(-CH₂-CO₋)$, 33.1, 30.8, 30.6, 30.5, 30.3, 26.0, 23.7 (-CH₂-lauroyl backbone), 14.5 $(CH_3$ -lauroyl). Note: ^{13}C NMR assignments marked with the same letter may be interchangeable.

Supplementary data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.tet.2005.10.046](http://dx.doi.org/doi:10.1016/j.tet.2005.10.046)

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Diastereoselective allylation of a-ketoamides bearing camphor N-tosylpyrazolidinone auxiliary: efficient synthesis of highly optically active two stereoisomers

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Abstract—Complementary allylation conditions were developed for the synthesis of both diastereomers of tertiary homoallylic alcohols. Treatment of camphor N-tosylpyrazolidinone derived a-ketoamides with allyltributylstannane afforded both the individual homoallylic alcohols in high optical purity (up to 98% de) when the reaction was carried out in the presence of $Sn(OTf)$ and $PdCl_2$, respectively. The stereochemical outcome and reversal of stereoselectivity in the reaction are proposed based on ¹³C NMR and FTIR studies. $©$ 2005 Elsevier Ltd. All rights reserved.

1. Introduction

The development of novel chiral auxiliaries that provide practical routes for the preparation of synthetically useful intermediates remains one of the most important fields in organic synthesis.^{[1](#page-121-0)} The fact that there are no universal chiral auxiliaries for all asymmetric transformations is a compelling reasons for discovering novel chiral auxiliaries for specific reactions.^{[2](#page-121-0)} From a practical synthetic point of view, the synthesis of both stereoisomers from the same chiral resources is attractive and has been a subject of considerable interest in recent years.^{[3](#page-121-0)} The reversal of stereochemistry from a single chiral starting material can be achieved by careful manipulation of the reaction components, especially the control of the architecture of the ligand-Lewis acid complex. The asymmetric allylation of ketones, 4 aldehydes^{[5](#page-121-0)} and imine^{[6](#page-122-0)} functionalities for the formation of homoallyl alcohols and homoallyl amines are well documented in the literature. The corresponding secondary and tertiary homoallylic alcohols are versatile intermediates that have been further used in organic synthesis.^{[7](#page-122-0)} However, allyl metal addition to α -keto esters and amides is a less well investigated topic.[8](#page-122-0) Diastereoselective allylation of camphor N -phenylpyrazolidinone derived α -ketoamides campnot *i* phenytpyrazon mode section ∞ accounting was examined recently by us.^{[9](#page-122-0)} The corresponding quarternary a-hydroxy carbonyls were obtained with good to excellent stereoselectivity. However, to the best of our knowledge, a diastereoselective allylation of α -ketoamides

with reversal of stereoselectivity has not been achieved to date. Here, we wish to report the asymmetric allylation of a novel camphor auxiliary derived α -ketoamides in the presence of a Lewis acid. The desired products were obtained in excellent diastereomeric excess in good to excellent chemical yields. The facial stereoselectivity and the stereochemical course of the reactions are discussed.

2. Results and discussion

The preparation of camphor N-tosylpyrazolidinone 4 followed the same synthetic route as described previously ([Scheme 1\)](#page-117-0).^{[2g](#page-121-0)} The treatment of $(+)$ -ketopinic acid 1 with p-toluenesulfonhydrazide producing the desired hydrazone 2 in nearly quantitative chemical yield. Cyclization was carried out by treatment with SOCl₂ in EtOAc at 75° C for 4 h. The corresponding C–N double bond was reduced with NaCNBH₃ in acetic acid to give the chiral camphor N-tosylpyrazolidinone 4 in 75% material yield. The desired α -ketoamides 5 are readily prepared by coupling the camphor N-tosylpyrazolidinone 4 with the freshly prepared a-ketocarboxylic acid chlorides at room temperature in 49–97% yields. It is noteworthy that both camphor N-tosylpyrazolidinone 4 and camphor N-tosylpyrazolidinone phenylglyoxylate 5a exist as two structural conformers in the asymmetric unit as indicated by X-ray crystallographic analyses.^{[10](#page-122-0)} Two distinct N-invertomers of camphor N-tosylpyrazolidinone 4 were observed in which the tosyl group is oriented toward and away from the C7 dimethyl group ([Scheme 1](#page-117-0), the dihedral angles of $NNSC_{SP}^2$ are -72.0 and $+84.1^{\circ}$, respectively).^{[11](#page-122-0)} On the other hand,

Keywords: Allylation; Reversal of stereoselectivity; Chiral auxiliary.

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Scheme 1. Reagents and conditions: (i) p-toluenesulfonhydrazide, CH_2Cl_2 , rt, 99%; (ii) SOCl₂, EtOAc, 75 °C, 90%; (iii) NaCNBH₃, AcOH, rt, 75%; (iv) RCOCOCl, $CH₂Cl₂$, base, rt, 49–97%.

the tosyl group points away from the C7 dimethyl group in 5a, thus avoiding the electrostatic repulsion with the dicarbonyl groups (the dihedral angles of $NNSC_{SP}^2$ are -69.9 and -83.0° , respectively). In addition, the amide carbonyl group in 5a is oriented away from the N-tosyl group with the dicarbonyl groups in an s-trans arrangement (the dihedral angles of COCO are 137.6 and 143.5° , respectively. See supporting information).

With the desired chiral α -ketoamides 5a–d in hand, we examined the diastereoselective allylation. The camphor N-tosylpyrazolidinone phenylglyoxylate 5a was chosen as a probe substrate. No reaction products were obtained when 5a was treated with allyltributylstannane in the absence of a Lewis acid. Various metal triflates such as $Sc(OTf)_{3}$, $Sm(OTf)_{3}$, $Zn(OTf)_{2}$ and $Eu(OTf)_{3}$ were systematically screened, but resulted in either low to moderate chemical yields or a low level of diastereoselectivities (data not shown). A satisfactory result was obtained when the reaction was carried out in the presence of 2.0 equiv of $Yb(OTf)$ ₃ for 24 h (Table 1, entry 1). The stereoselectivity was further improved to an excellent level when $Sn(OTf)_{2}$ was employed in a 5 min reaction (entry 2). The diastereoselectivity was determined by H NMR and HPLC studies of relevant peaks. The absolute stereochemistry of the newly generated stereogenic center in the major diasteoreomer $6a$ was assigned to have an R configuration, deduced from the single crystal X-ray analysis. Interestingly, the sense of stereoinduction was reversed with less reactivity when the reaction was carried out in the presence of $PdCl₂$ and the structure of product $7a$ was again confirmed by single crystal X-ray analysis (entry 3). A careful examination of the ¹H NMR spectra indicates that the characteristic C-2 methine proton (camphor numbering) appears at 3.16 ppm for the R isomer while at 4.32 ppm for the counter \overrightarrow{S} isomer.^{[9,12](#page-122-0)} The diamagnetic anisotropy effect of the aromatic substituent may account for the shielding effect.

Having identified two discrete Lewis acids $[Sn(OTf)]$ ₂ and PdCl₂] that can produce complementary diastereomers of the allylation reactions, we sought to examine the nature of the architecture of the chiral auxiliary backbone. Toward this end, the allylation proceeded smoothly when camphor N-tosylpyrazolidinone derived 2-thienylglyoxylate 5b was used under the optimum reaction conditions, providing 6b with excellent selectivity in the presence of $Sn(OTf)₂$ (entry 4). The opposite diasteoreomer with a similar selectivity was produced when $PdCl₂$ was used (entry 5).

Table 1. Diastereoselective allylation of camphor N-tosylpyrazolidinone derived α -ketoamides 5a–d in the presence of a Lewis acid^a

^a Unless otherwise specified, all reactions were carried out in a solvent [in CH₃CN when Sn(OTf)₂ was used and in CH₂Cl₂ when PdCl₂ was used] at room temperature using 5 (0.11 mmol), allyltributylstannane (1.0

 \degree Total isolated yield (6+7).
 \degree The absolute stereochemistry of the newly generated stereogenic center was deduced by X-ray crystallographic analyses.

 $\rm ^d$ Ratios of diastereomers were determined by $\rm ^1H$ NMR analysis of relevant peaks and HPLC analyses of crude products.

 $^{\circ}$ The deacylated compound N-tosylcamphorpyrazolidinone-3-indoloylglyoxylate was isolated in 50%. $^{\circ}$ Absolute stereochemistry are assigned by analogy.

It is noteworthy that, under either reaction conditions, no desired allylation product was observed with camphor N-tosylpyrazolidinone derived 3-indoloylglyoxylate. This was presumably due to the presence of an active hydrogen atom that decreases the reactivity. This problem was eliminated when NH protected analogue was used. Thus, camphor N-tosylpyrazolidinone-1-acetyl-3-indoloylglyoxylate 5c upon allylation afforded the expected product with fairly high yield and selectivity in the presence of $Sn(OTf)$ ₂ (entry 6). However, no desired reaction products were observed when $PdCl₂$ was employed and the deacylation product was isolated (entry 7). Finally, the aliphatic substrate 5d efficiently participates in the reaction, leading to the desired products with good diastereoselectivity, but the reactivity decreased dramatically when $PdCl₂$ was used as a Lewis acid (entries 8–9).

The high degree of stereoselectivity and the reversal of stereoselectivity can be rationalized by the chelation and non-chelation control of the Lewis acid with α -ketoamides 5 as shown in Figure 1. Among the possible conformations, the pseudo planar s-trans conformer of the α -dicarbonyl group in 5 is electronically favored over its s-cis conformer to avoid electrostatic repulsive interactions between the adjacent carbonyl groups. This was confirmed by X-ray crystallographic analyses of 5a,b in the solid states. The reversal of stereoselectivity can be explained by simple 13 C NMR and FTIR studies. When a mixture of $Sn(OTf)$ ₂ and 5a in CD₃CN was examined by 13 C NMR, no significant chemical shift difference in the carbonyl region was observed $(<0.05$ ppm) in comparison with a spectrum of the pure substrate.[13](#page-122-0) This indicates that there is little chelation of Lewis acid metal atom with the carbonyl oxygen atoms. In addition, no significant carbonyl group stretching band changes in the infrared spectrum of the mixture were observed. However, a new stretching band corresponding to an S=O group developed at 1272 cm^{-1} , which is different from the stretching band of 1240 cm^{-1} in 5a.^{[14](#page-122-0)} On the other hand, a mixture of $PdCl_2$ and substrate 5a give no significant carbonyl signal shift difference or absorption band changes (carbonyl and sulfonyl groups) by 13 C NMR and FTIR spectroscopy. These data indicate a strong coordination of the $Sn(OTf)$, with sulfonyl oxygen atoms and this may account for the much faster reaction in the presence of $Sn(OTf)_2$ in comparison to the use of $PdCl_2$ (5 min vs 24 h). The coordination of the tin to the sulfonyl group resulted in the rapid release of the nucleophilic allyl group. The allyl group then attacks the α -carbonyl amide group from the bottom si face to afford the desired product.

Figure 1. Proposed mechanistic explanation of the diastereoselective allylation. The stereoview of the Chem3D structure of 5a was generated from the X-ray crystal coordinates. All hydrogen atoms are omitted for the sake of clarity.

While, in the case of $PdCl₂$, the electronic rich tosyl group prevents nucleophilic addition from the si face, leading to the addition from the top re face.^{[15](#page-122-0)}

3. Conclusion

In conclusion, a new camphor-based chiral auxiliary N-tosylpyrazolidinone was synthesized, and tested as a stereocontrolling element in allylation reactions across a range of substrates. The present work confirms that the correct choice of Lewis acid is essential in determining the diastereoselective outcome of the allylation.^{[16](#page-122-0)} The strongly chelating $Sn(OTf)_2$ led to the si face addition diastereomer, while the non-chelating $PdCl₂$ gave the counter diastereomer with allyltributyltin. Further synthetic applications are currently underway.

4. Experimental

4.1. General methods

All reagents were used as purchased from the commercial suppliers without further purification. NMR spectra were recorded on a Bruker Avance 400 NMR spectrometer (400 MHz for ${}^{1}H$, and 100 MHz for ${}^{13}C$). Chemical shifts are reported in δ ppm referenced to an internal TMS standard for ¹H NMR and chloroform-d (δ 77.0) for ¹³C NMR. Optical rotations were measured on a JASCO P-1010 polarimeter. Infrared spectra were recorded on a Perkin-Elmer FTIR spectrum 500 spectrometer. Only the characteristic peaks are quoted. EI mass spectra were recorded on Finnigan TSQ-700 at an ionizing energy of 70 eV and HRMS spectra were recorded on JEOL SX-102A. Routine monitoring of reactions was performed using silica gel, glass-backed TLC plates (Merck Kieselgel 60 F254) and visualized by UV light (254 nm). Analytical highperformance liquid chromatography was performed on a JASCO PU-1580 HPLC (ZORBAX analytical NH₂ column). Solutions were evaporated to dryness under reduced pressure with a rotary evaporator and the residue was purified by flash column chromatography on silica gel (230–400 mesh) with the indicated eluents. Air and/or moisture sensitive reactions were performed under usual inert atmosphere conditions.

Crystallographic data for the structures in this paper have been deposited at the Cambridge Crystallographic Data Center and allocated the deposit numbers CCDC 281384-281390, and 284467.[17](#page-122-0)

4.1.1. 2-(2-Tosylhydrazono)-7,7-dimethylbicyclo[2.2.1] heptane-1-carboxylic acid (2). To a solution of $(+)$ ketopinic acid 1 (20 g, 109.9 mmol) in CH_2Cl_2 (300 mL) was added p-toluenesulfonhydrazide (22.5 g, 120.9 mmol) in one portion and the mixture was allowed to stir for 4 h at room temperature. The reaction mixture was diluted with water (500 mL) and extracted with dichloromethane $(3 \times 500 \text{ mL})$, the organic layer was separated and dried over anhydrous $MgSO₄$ and concentrated to give ketopinic acid tosylhydrazone 2 as a white solid (38.1 g; 99%); mp: 114–115 °C. R_f = 0.29 (1:1 hexanes/EtOAc). $[\alpha]_D^{27}$ +47.1

(c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 7.81 (d, 2H, $J=7.6$ Hz), 7.74 (br s, 1H, –NH–, D₂O exchangeable), 7.35 (d, 2H, $J=7.6$ Hz), 2.51–2.39 (m, 2H), 2.43 (s, 3H), $2.09-2.03$ (m, 2H), 1.94 (d, 1H, $J=17.4$ Hz), 1.68–1.62 (m, 1H), 1.33–1.28 (m, 1H), 1.20 (s, 3H), 0.79 (s, 3H); 13C NMR (100 MHz, CDCl3) d 171.9, 169.0, 145.0, 134.3, 130.1, 127.9, 61.1, 51.7, 44.3, 33.9, 31.1, 27.5, 21.5, 19.7, 18.3; IR $(\text{neat}, \text{ cm}^{-1})$: 3436–2408 (br), 1715, 1597, 1416, 1337, 1166, 1088, 920; HRMS (EI) calcd for $C_{17}H_{22}N_2O_4S$ 350.1295. Found 350.1283.

4.1.2. 10,10-Dimethyl-3-N-tosyl-4-aza-tricyclo $[5.2.1.0^{1.5}]$ dec-4-en-2-one (3). The above prepared ketopinic acid tosylhydrazone 2 (20 g, 57.14 mmol) was dissolved in ethyl acetate (300 mL). To this solution was added thionyl chloride (15.8 mL, 217.14 mmol) slowly and the reaction mixture was brought to reflux at 75 °C for 4 h under N_2 atmosphere. The reaction mixture quenched with water (600 mL) and extracted with ethyl acetate $(3 \times 600 \text{ mL})$. The dried $MgSO₄$ extract was concentrated in vacuo and purified by flash column chromatography, eluted with (1:1 hexanes/EtOAc) afforded the cyclized product 3 as a colorless solid (17.07 g; 90%).; mp: 83–84 °C. R_f =0.44 (1:1 hexanes/EtOAc). $[\alpha]_D^{27}$ – 99.6 (c 1, CHCl₃); ¹H NMR $(400 \text{ MHz}, \text{CDCl}_3)$ δ 7.93 (d, 2H, J=7.9 Hz), 7.32 (d, 2H, $J=7.9$ Hz), 2.59 (d, 1H, $J=18.0$ Hz), 2.43 (s, 3H), 2.22–2.20 (m, 2H), 2.15 (d, 1H, $J=18.0$ Hz), 2.10–2.05 (m, 1H), 1.76–1.69 (m, 1H), 1.52–1.45 (m, 1H), 1.12 (s, 3H), 0.63 (s, 3H); ¹³C NMR (100 MHz, CDCl₃) δ 176.2, 172.2, 145.2, 135.3, 129.7, 127.9, 64.0, 50.9, 48.7, 32.2, 26.8, 26.0, 21.5, 18.5, 18.1; IR (neat, cm⁻¹): 3065, 2961, 2923, 2853, 1760, 1747, 1645, 1598, 1455, 1373, 1295, 1258, 1178, 1082, 959, 814, 705; HRMS (EI) calcd for $C_{17}H_{20}N_2O_3S$ 332.1189. Found 332.1190; Crystal data for 3 (colorless crystal, recrystallized from hexanes/EtOAc) at 25 °C: C₁₇H₂₀N₂O₃S, *M* 332.422, Orthorhombic, $P2_12_12_1$, $a=10.1101(2)$ \AA , $b=12.9076(3)$ \AA , $c=25.9674(8)$ \AA , $V=$ 3388.67 (15) \AA^3 , Z=8, D_x=1.303 Mg/m³, μ =0.21 mm⁻¹, 2336 reflections, 416 parameters, $R = 0.058$, $R_w = 0.096$.

4.1.3. 10,10-Dimethyl-3-N-tosyl-4-aza-tricyclo $[5.2.1.0^{1,5}]$ decan-2-one (4). To a stirred solution of cyclized substrate 3 $(5.05 \text{ g}, 15.21 \text{ mmol})$ in AcOH (75 mL) at room temperature was added slowly NaBH3CN (13.19 g, 210 mmol). The reaction mixture was stirred at room temperature for 30 min until the reaction was completed and diluted with H_2O (750 mL) and extracted with CH_2Cl_2 (3×600 mL). The organic layer was dried over anhydrous $MgSO₄$ and concentrated to give a crude product, which is purified by flash column chromatography eluted with $(2:1$ hexanes/ EtOAc) to yield 4 (3.75 g; 75%) as a white solid; mp: 118–119 °C. R_f = 0.38 (2:1 hexanes/EtOAc). [α] $^{27}_{D}$ + 14.4 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 7.93 (d, 2H, J= 8.0 Hz), 7.32 (d, 2H, $J=8.0$ Hz), 5.05 (br s, 1H, –NH–), 3.5 (dd, 1H, $J=8.4$, 4.0 Hz), 2.44 (s, 3H), 2.11–2.02 (m, 2H), $1.89-1.80$ (m, 2H), 1.66 (dd, 1H, $J=13.2$, 8.4 Hz), 1.26–1.15 (m, 2H), 0.94 (s, 3H), 0.76 (s, 3H); 13C NMR (100 MHz, CDCl3) d 172.3, 145.2, 134.5, 129.4, 128.4, 64.5, 58.9, 52.5, 47.2, 36.4, 29.0, 26.4, 21.6, 20.4, 19.7; IR $(neat, cm⁻¹)$: 3323, 3302, 2958, 2899, 1757, 1747, 1598, 1481, 1360, 1173, 1075, 863, 817; HRMS (EI) calcd for $C_{17}H_{22}N_2O_3S$ 334.1346. Found 334.1350. Anal. Calcd for $C_{17}H_{22}N_{2}O_{3}S$: C, 61.05; H, 6.63; N, 8.38. Found: C, 61.07;

H, 6.60; N, 8.20; Crystal data for 4 (colorless crystal, recrystallized from hexanes/EtOAc) at 25° C: C₁₇H₂₂N₂O₃S, M 334.438, Monoclinic, $P2_1$, $a=9.7297(3)$ Å, $b=$ 14.6999(4) Å, $c = 12.2705(5)$ Å, $V = 1714.77(10)$ Å³, $Z = 4$, $D_x = 1.295$ Mg/m³, $\mu = 0.205$ mm⁻¹, 5864 reflections, 415 parameters, $R=0.0643$, $R_w=0.1547$.

4.2. General procedure for the synthesis of compounds 5a–c

Under N_2 atmosphere a mixture of benzoylformic acid (2.25 g, 15.0 mmol) and thionyl chloride (15.27 mL, 209.29 mmol) was refluxed at 75° C for 2 h and concentrated on rotary evaporator to remove $S OCl₂$. The resulting crude mixture was dissolved in CH_2Cl_2 (15 mL) and added to a solution of camphor N-tosylpyrazolidinone $4(1.0 \text{ g})$, 2.99 mmol) in CH_2Cl_2 (10 mL). The reaction mixture was allowed to stir for 15 min and quenched with water (50 mL) and extracted with CH_2Cl_2 (3×75 mL). The organic extracts were combined, dried over anhydrous $MgSO₄$ and concentrated to give crude products, which were subject to flash column chromatography eluted with $(2:1$ hexanes/ EtOAc) afforded the pure product $5a(1.35 g, 97\%)$ as a white solid.

4.2.1. 10,10-Dimethyl-3-N-tosyl-4-N-(2-oxo-2-phenylacetyl)-tricyclo[5.2.1.0^{1,5}]decan-2-one (5a). $R_f = 0.38$ (2:1) hexanes/EtOAc); mp: 179–181 °C. $[\alpha]_D^{27}$ – 194.6 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 8.15 (d, 2H, J= 7.6 Hz), 7.92 (s, 2H), 7.65 (t, 1H, $J=7.2$ Hz), 7.53 (t, 2H, $J=7.6$ Hz), 7.37 (d, 2H, $J=8.4$ Hz), 4.07 (s, 1H), 2.46 (s, 3H), 2.11 (t, 1H, $J=9.8$ Hz), 1.95–1.90 (m, 2H), 1.77 (dd, 1H, $J=13.4$, 8.2 Hz), 1.26 (s, 3H), 1.26–1.22 (m, 2H), 1.08 (m, 1H), 1.08 (s, 3H); ¹³C NMR (100 MHz, CDCl₃) δ 171.3, 162.2, 146.0, 134.4, 133.6, 133.0, 130.5, 130.3, 130.0, 128.6, 128.4, 70.0, 60.0, 52.8, 47.5, 36.1, 28.8, 26.5, 21.6, 20.1, 20.0; IR (neat, cm⁻¹): 3065, 3003, 2964, 2925, 1768, 1680, 1652, 1594, 1483, 1450, 1385, 1241, 1173, 1083, 946, 816, 732; HRMS (EI) calcd for $C_{25}H_{26}N_{2}O_{5}S$ 466.1557. Found 466.1574. Anal. Calcd for C₂₅H₂₆N₂O₅S: C, 64.36; H, 5.62; N, 6.00. Found: C, 64.33; H, 5.62; N, 5.75; Crystal data for 5a (colorless crystal, recrystallized from hexanes/EtOAc) at 20 °C: C_2 ₅H₂₆N₂O₅S, *M* 466.54, Monoclinic, *P*₂₁, *a* = 10.1710(5) A, $b=11.8910(5)$ A, $c=19.5700(12)$ A, $V=$ 2360.8(2) \mathring{A}^3 , Z=4, D_x=1.313 Mg/m³, μ =0.176 mm⁻¹, 7815 reflections, 596 parameters, $R=0.1298$, $R_w=0.1964$.

4.2.2. 10,10-Dimethyl-3-N-tosyl-4-N-[2-oxo-2-(thiophen-2-yl)acetyl]-tricyclo^{[5.2.1.0^{1,5}]decan-2-one (5b). White} solid. R_f =0.35 (2:1 hexanes/EtOAc); mp: 180–182 °C. $[\alpha]_D^{27}$ – 232.7 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 8.13 (d, 1H, $J=3.6$ Hz), 7.93 (s, 2H), 7.81 (d, 1H, $J=$ 4.4 Hz), 7.37 (d, 2H, $J=8.0$ Hz), 7.2 (t, 1H, $J=4.4$ Hz), 4.11 (d, 1H, $J=6.8$ Hz), 2.46 (s, 3H), 2.09–2.06 (m, 1H), $1.94-1.92$ (m, 2H), 1.82 (dd, 1H, $J=14.0$, 8.4 Hz), 1.26–1.22 (m, 2H), 1.18 (s, 3H), 1.05 (m, 1H), 1.05 (s, 3H); ¹³C NMR (100 MHz, CDCl₃) δ 179.1, 174.3, 161.4, 146.0, 139.5, 137.0, 134.6, 129.8, 129.1, 128.9, 128.7, 70.5, 60.3, 52.6, 47.4, 36.8, 28.8, 26.7, 21.8, 20.3, 20.0; IR (neat, cm^{-1}): 3106, 2997, 2962, 2920, 2884, 1766, 1652, 1596, 1409, 1380, 1357, 1246, 1175, 1083, 1055, 815; HRMS (EI) calcd for $C_{23}H_{24}N_2O_5S_2$ 472.1127. Found 472.1129. Anal. Calcd for $C_{23}H_{24}N_2O_5S_2$: C, 58.45; H, 5.12; N, 5.93. Found: C, 58.68; H, 5.11; N, 5.88; Crystal data for 5b (colorless crystal, recrystallized from hexanes/EtOAc) at 20° C: $C_{23}H_{24}N_2O_5S_2$, *M* 472.56, Monoclinic, *P*2₁, *a*= 12.5900(4) A^{h} , $b=6.6550(2)$ A^{h} , $c=13.2260(5)$ A^{h} , $V=$ 1107.62(6) \mathring{A}^3 , Z=2, D_x=1.417 Mg/m³, μ =0.279 mm⁻¹, 3697 reflections, 285 parameters, $R = 0.0735$, $R_w = 0.1837$.

4.2.3. 10,10-Dimethyl-3-N-tosyl-4-N-[2-(1-acetyl-1Hindol-3-yl)-2-oxoacetyl]-tricyclo $[5.2.1.0^{1.5}]$ decan-2-one (5c). White solid. $R_f = 0.59$ (2:1 hexanes/EtOAc); mp: 194–195 °C; ¹H NMR (400 MHz, CDCl₃) δ 8.69 (s, 1H), 8.49–8.47 (m, 1H), 8.36–8.34 (m, 1H), 7.94 (s, 2H), 7.48–7.42 (m, 2H), 7.36 (d, 2H, $J=7.6$ Hz), 4.22 (s, 1H), 2.75 (s, 3H), 2.46 (s, 3H), 2.12 (td, 1H, $J=11.7$, 3.9 Hz), 1.96–1.93 (m, 2H), 1.86 (dd, 1H, $J=13.7$, 8.0 Hz), 1.31–1.24 (m, 3H), 1.21 (s, 3H), 1.07 (s, 3H); 13C NMR (100 MHz, CDCl3) d 180.0, 169.0, 162.2, 146.1, 136.2, 135.9, 129.8, 129.0, 127.1, 126.7, 125.6, 122.0, 117.7, 116.6, 70.7, 60.4, 52.7, 47.6, 29.7, 28.9, 26.7, 23.9, 21.8, 20.3, 20.1; IR (neat, cm⁻¹): 3059, 2962, 2925, 1768, 1732, 1652, 1539, 1448, 1379, 1212, 1175, 1008, 756; HRMS (EI) calcd for $C_{29}H_{29}N_3O_6S$ 547.1777. Found 547.1779.

4.2.4. 10,10-Dimethyl-3-N-tosyl-4-N-(2-oxobutanoyl) tricyclo[5.2.1.0^{1,5}]decan-2-one (5d). Under N_2 atmosphere a mixture of 2-ketobutyric acid (3.00 g, 29.30 mmol) and thionyl chloride (32.1 mL, 440.7 mmol) was refluxed at 75° C for 2 h and concentrated on rotary evaporator to remove SOCl₂. The resulting crude product was dissolved in CH_2Cl_2 (5 mL) and added to a solution of camphor N -tosylpyrazolidinone 4 (0.20 g, 0.59 mmol) in CH₂Cl₂ (5 mL). This was added pyridine (2.30 mL, 29.50 mmol) and the reaction was allowed to stir at room temperature for 12 h. The reaction mixture was then quenched with H_2O (20 mL) and extracted with CH_2Cl_2 (3 × 20 mL). The combined organic extracts dried over anhydrous MgSO₄ and the crude products were purified by flash column chromatography eluted with (2:1 hexane/EtOAc) provided product 5d (0.12 g; 50%) as a colorless oil. $R_f = 0.38$ (2:1) hexanes/EtOAc); ¹H NMR (400 MHz, CDCl₃) δ 7.81 (d, 2H, $J=8.2$ Hz), 7.40 (d, 2H, $J=8.2$ Hz), 3.78 (m, 1H), 3.11 (dq, 1H, $J=19.0$, 7.2 Hz), 2.81 (dq, 1H, $J=19.0$, 7.2 Hz), 2.48 (s, 3H), 2.03–1.86 (m, 4H), 1.81 (dd, 1H, $J=14.3$, 8.2 Hz), $1.26-1.20$ (m, 2H), 1.16 (t, 3H, $J=7.2$ Hz), 1.16 (s, 3H), 1.03 (s, 3H); ¹³C NMR (100 MHz, CDCl₃) δ 197.0, 174.9, 162.3, 146.4, 133.9, 130.0, 128.6, 70.3, 60.0, 52.4, 47.3, 35.2, 31.7, 28.7, 26.6, 21.6, 20.2, 19.8, 6.9; IR (neat, cm⁻¹): 2964, 2941, 2889, 1769, 1726, 1660, 1596, 1458, 1379, 1189, 1175, 1086, 964, 815; HRMS (EI) calcd for $C_{21}H_{26}N_{2}O_{5}S$ 418.1562. Found 418.1560.

4.3. Typical procedure for the synthesis of compounds 6a and 7a

To a solution of camphor N-tosylpyrazolidinone derived α -ketoamide (51.27 mg, 0.11 mmol) in CH₃CN (4 mL) was added allyltributylstannane (0.37 mL, 1.06 mmol) and tin(II) triflate $(91.7 \text{ mg}, 0.22 \text{ mmol})$. The reaction was allowed to stir for 5 min at room temperature. The mixture was quenched with H_2O (15 mL) and extracted with CH_2Cl_2 $(3 \times 20 \text{ mL})$. The organic layers were separated and dried over anhydrous $MgSO₄$ and concentrated to give crude

products. Further purificatin by flash column chromatography eluted with (3:1 hexanes/EtOAc) afforded 6a and 7a as a white solid (51.42 mg; 92%, ratio of diastereomers 99:01).

4.3.1. 10,10-Dimethyl-3-N-tosyl-4-N-[(R)-2-hydroxy-2-phenylpent-4-enoyl]-tricyclo^{[5.2.1.0^{1,5}]decan-2-one} (6a). $R_f = 0.54$ (2:1 hexanes/EtOAc); mp: 170–172 °C. $[\alpha]_D^{27}$ $+$ 142.0 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 8.14 (d, 2H, $J=8.0$ Hz), 7.54 (d, 2H, $J=7.2$ Hz), 7.37 (d, 2H, $J=8.0$ Hz), 7.33–7.26 (m, 3H), 5.92 (m, 1H), 5.35 (d, 1H, $J=9.6$ Hz), 5.28 (d, 1H, $J=16.8$ Hz), 3.34 (dd, 1H, $J=$ 13.2, 5.5 Hz), 3.16 (s, 1H), 3.02 (s, 1H), 2.76–2.73 (m, 1H), 2.46 (s, 3H), 2.46–2.42 (m, 1H), 2.00–1.94 (m, 1H), 1.81– 1.78 (m, 3H), 1.27 (s, 3H), 1.05 (s, 3H), 0.97–0.91 (m, 1H), 0.86–0.81 (m, 1H); ¹³C NMR (100 MHz, CDCl₃) δ 171.7, 153.0, 144.8, 139.5, 136.0, 132.6, 129.3, 129.1, 128.4, 128.0, 124.7, 122.1, 78.4, 67.5, 59.0, 55.1, 49.5, 46.3, 41.5, 28.0, 26.2, 21.7, 20.7, 19.7; IR (neat, cm⁻¹): 3530, 3065, 2965, 2884, 1748, 1703, 1450, 1368, 1173, 1083, 814, 739; HRMS (EI) calcd for $C_{28}H_{32}N_2O_5S$ 509.2060. Found 509.2053; Crystal data for 6a (colorless crystal, recrystallized from hexanes/EtOAc) at 20 °C: $C_{28}H_{32}N_2O_5S$, M 508.62, Monoclinic, $P2_{1}$, $a=8.6460(2)$ Å, $b=$ 11.5040(3) Å, $c = 13.4890(3)$ Å, $V = 1301.10(5)$ Å³, $Z = 2$, $D_x = 1.298$ Mg/m³, $\mu = 0.165$ mm⁻¹, 4476 reflections, 326 parameters, $R=0.0652$, $R_w=0.1377$.

4.3.2. 10,10-Dimethyl-3-N-tosyl-4-N-[(S)-2-hydroxy-2-phenylpent-4-enoyl]-tricyclo[5.2.1.0^{1,5}]decan-2-one (7a). White solid. $R_f = 0.46$ (2:1 hexanes/EtOAc); mp: 194–195 °C. $[\alpha]_D^{27}$ – 71.3 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl₃) δ 8.08 (d, 2H, J=7.8 Hz), 7.67 (d, 2H, J=7.8 Hz), 7.39–7.28 (m, 5H), 5.94 (m, 1H), 5.32 (d, 1H, $J=10.1$ Hz), 5.26 (d, 1H, $J=17.2$ Hz), 4.32 (t, 1H, $J=6.2$ Hz), 3.27 (dd, 1H, $J=13.4$, 6.2 Hz), 2.93 (br s, 1H), 2.66 (dd, 1H, $J=13.4$, 8.4 Hz), 2.46 (s, 3H), 2.08–2.02 (m, 1H), 1.89–1.74 (m, 3H), 1.56 (s, 1H), 1.27–1.16 (m, 2H), 0.90 (s, 3H), 0.70 (s, 3H); 13C NMR (100 MHz, CDCl₃) δ 174.0, 172.9, 145.2, 140.2, 135.6, 133.0, 129.5, 128.9, 128.1, 127.6, 125.4, 121.9, 78.7, 72.5, 60.3, 51.7, 51.1, 47.0, 39.5, 29.1, 26.5, 21.8, 20.2, 19.5; IR $(neat, cm⁻¹)$: 3504, 3065, 2965, 2935, 1755, 1668, 1447, 1379, 1295, 1189, 1174, 1086; HRMS (EI) calcd for $C_{28}H_{32}N_2O_5S$ 508.2026. Found 508.2009; Crystal data for 7a (colorless crystal, recrystallized from hexanes/EtOAc) at -73 °C: C₂₈H₃₂N₂O₅S, *M* 508.62, Orthorhombic, *P*2₁2₁2₁, $a=6.3790(2)$ Å, $b=18.5760(6)$ Å, $c=21.1690(8)$ Å, $V=$ 2508.45(15) \hat{A}^3 , Z=4, D_x=1.347 Mg/m³, μ =0.172 mm⁻¹, 4409 reflections, 326 parameters, $R=0.1054$, $R_w=0.1766$.

4.3.3. 10,10-Dimethyl-3-N-tosyl-4-N-[(S)-2-hydroxy-2- $(thiophen-2-yl)$ pent-4-enoyl]-tricyclo[5.2.1.0^{1,5}] decan-2one (6b). White solid. $R_f = 0.51$ (2:1 hexanes/EtOAc); mp: 163–165 °C. $[\alpha]_D^{27}$ + 122.8 (c 1, CHCl₃); ¹H NMR $(400 \text{ MHz}, \text{CDCl}_3)$ δ 8.11 (d, 2H, $J=8.0 \text{ Hz}$), 7.35 (d, 2H, $J=8.0$ Hz), 7.20 (d, 1H, $J=4.8$ Hz), 7.01 (d, 1H, $J=$ 3.7 Hz), 6.92 (t, 1H, $J=4.8$ Hz), 5.89 (m, 1H), 5.36 (d, 1H, $J=10.1$ Hz), 5.31 (d, 1H, $J=17.2$ Hz), 3.52 (s, 1H), 3.35 (dd, 1H, $J=13.3$, 5.1 Hz), 3.25 (br s, 1H), 2.72–2.65 (m, 2H), 2.44 (s, 3H), 2.06–1.98 (m, 2H), 1.87–1.81 (m, 2H), 1.25 (s, 3H), 1.12–1.06 (m, 1H), 1.06 (s, 3H), 0.97 (t, 1H, $J=10.0$ Hz); ¹³C NMR (100 MHz, CDCl₃) δ 177.6, 171.8, 144.9, 144.3, 135.8, 132.0, 129.2, 129.1, 127.3, 125.0,

124.2, 122.4, 78.0, 68.2, 59.0, 55.1, 49.7, 46.3, 41.7, 28.0, 26.2, 21.7, 20.6, 19.7; IR (neat, cm⁻¹): 3516, 2992, 2966, 2884, 1748, 1698, 1452, 1368, 1207, 1173, 1086, 815; HRMS (EI) calcd for $C_{26}H_{30}N_2O_5S_2$ 514.1591. Found 514.1599; Crystal data for 6b (colorless crystal, recrystallized from hexanes/EtOAc) at -123 °C: C₂₆H₃₀N₂O₅S₂, M 514.64, Monoclinic, $P2_1$, $a=8.5300(3)$ Å, $b=$ 11.2470(4) Å, $c = 13.4740(7)$ Å, $V = 1247.57(9)$ Å³, $Z = 2$, $D_x = 1.370$ Mg/m³, $\mu = 0.254$ mm⁻¹, 4278 reflections, 317 parameters, $R=0.0625$, $R_w=0.1386$.

4.3.4. 10,10-Dimethyl-3-N-tosyl-4-N-[(R)-2-hydroxy-2- $(thiophen-2-yl)$ pent-4-enoyl]-tricyclo $[5.2.1.0^{1,5}]$ decan-**2-one (7b).** White solid. $R_f=0.41$ (2:1 hexanes/EtOAc); mp: 167–168 °C. $[\alpha]_D^{27}$ – 50.1 (c 1, CHCl₃); ¹H NMR $(400 \text{ MHz}, \text{CDCl}_3)$ δ 8.06 (d, 2H, $J=8.2 \text{ Hz}$), 7.37 (d, 2H, $J=8.2$ Hz), 7.25 (dd, 1H, $J=5.0$, 1.0 Hz), 7.17 (dd, 1H, $J=$ 3.5, 1.0 Hz), 6.95 (dd, 1H, $J=5.0$, 3.5 Hz), 5.95 (m, 1H), 5.34 (d, 1H, $J=10.8$ Hz), 5.30 (d, 1H, $J=18.0$ Hz), 4.34 (s, 1H), 3.25 (dd, 1H, $J=13.6$, 7.5 Hz), 3.18 (br s, 1H), 2.89 (dd, 1H, $J=13.6$, 7.5 Hz), 2.46 (s, 3H), 2.19–2.16 (m, 1H), 2.09–2.04 (m, 1H), 1.92 (dd, 1H, $J=14.4$, 8.1 Hz), $1.87-1.81$ (m, 1H), 1.67 (t, 1H, $J=3.7$ Hz), 1.30-1.19 (m, 2H), 0.94 (s, 3H), 0.81 (s, 3H); 13C NMR (100 MHz, CDCl3) d 174.0, 172.8, 145.3, 144.5, 135.4, 132.4, 129.5, 128.9, 126.8, 125.3, 124.4, 122.3, 78.3, 72.8, 60.4, 51.7, 51.5, 47.0, 39.9, 29.1, 26.6, 21.8, 20.3, 19.5; IR (neat, cm⁻¹): 3501, 3054, 2992, 2961, 1741, 1683, 1380, 1175, 1080, 1039, 936, 827, 752; HRMS (EI) calcd for $C_{26}H_{30}N_2O_5S_2$ 514.1591. Found 514.1598. Anal. Calcd for $C_{26}H_{30}N_2O_5S_2$: C, 60.68; H, 5.88; N, 5.44. Found: C, 60.82; H, 5.85; N, 5.45; Crystal data for 7b (colorless crystal, recrystallized from hexanes/EtOAc) at -73 °C: $C_{26}H_{30}N_2O_5S_2$, *M* 514.64, Orthorhombic, $P_212_12_1$, $a=$ 9.8170(2) Å, $b=12.1720(2)$ Å, $c=21.7010(5)$ Å, $V=$ 2593.11(9) \mathring{A}^3 , Z=4, D_x=1.318 Mg/m³, μ =0.244 mm⁻¹, 4469 reflections, 316 parameters, $R=0.0531$, $R_w=0.1272$.

4.3.5. 10,10-Dimethyl-3-N-tosyl-4-N-[(R)-2-(1-acetyl-1Hindol-3-yl)-2-hydroxypent-4-enoyl]-tricyclo $[5.2.1.0^{1,5}]$ decan-2-one (6c). White solid. $R_f=0.60$ (2:1 hexanes/ EtOAc); mp: 153–155 °C. $[\alpha]_D^{16}$ + 157.3 (c 1, CHCl₃); ¹H NMR (400 MHz, CDCl3) δ 8.44 (d, 1H, J = 7.9 Hz), 8.10 (d, 2H, $J=8.2$ Hz), 7.96 (d, 1H, $J=7.9$ Hz), 7.49 (s, 1H), 7.38 (d, 2H, $J=8.2$ Hz), 7.35 (t, 1H, $J=7.9$ Hz), 7.26 (t, 1H, $J=$ 7.9 Hz), 5.94 (m, 1H), 5.35 (d, 1H, $J=10.4$ Hz), 5.30 (d, 1H, $J=17.4$ Hz), 3.54 (dd, 1H, $J=7.7$, 5.1 Hz), 3.43 (dd, 1H, $J=12.6$, 4.1 Hz), 3.25 (br s, 1H), 2.80–2.75 (m, 1H), 2.69 (d, 1H, $J=12.6$ Hz), 2.61 (s, 3H), 2.47 (s, 3H), 1.98 (td, 1H, $J=12.0$, 4.8 Hz), 1.83–1.78 (m, 3H), 1.30–1.26 (m, 1H), 1.16 (s, 3H), 1.03 (s, 3H), 1.00–0.96 (m, 1H); 13C NMR (100 MHz, CDCl3) d 176.0, 172.2, 168.6, 145.2, 136.4, 135.5, 132.0, 129.3, 129.2, 127.3, 125.5, 123.8, 122.9, 122.1, 121.9, 121.7, 116.5, 77.4, 59.2, 47.4, 46.4, 41.2, 31.5, 28.1, 26.3, 22.6, 21.7, 20.5, 19.7, 14.1; IR (neat, cm⁻¹): 3418, 3080, 2962, 2930, 2894, 1748, 1710, 197, 1451, 1382, 1331, 1225, 1173, 1083, 815; HRMS (EI) calcd for $C_{32}H_{35}N_3O_6S$ 589.2247. Found 589.2240.

4.3.6. 10,10-Dimethyl-3-N-tosyl-4-N-[(S)-2-ethyl-2 hydroxypent-4-enoyl]-tricyclo[5.2.1.0^{1,5}]decan-2-one (6d). White solid. $R_f = 0.42$ (2:1 hexanes/EtOAc); mp: 138–140 °C; ¹H NMR (400 MHz, CDCl₃) δ 8.10 (d, 2H, J =

8.2 Hz), 7.35 (d, 2H, $J=8.2$ Hz), 6.05 (m, 1H), 5.34 (d, 1H, $J=10.0$ Hz), 5.24 (d, 1H, $J=17.1$ Hz), 3.90 (s, 1H), 2.87 (dd, 1H, $J=13.3$, 5.2 Hz), 2.67–2.58 (m, 1H), 2.44 (s, 3H), 2.38–2.28 (m, 1H), 2.22–2.11 (m, 2H), 1.98–1.87 (m, 2H), 1.70–1.65 (m, 1H), 1.34 (s, 3H), 1.30–1.23 (m, 3H), 1.13 (s, 3H), 0.93 (t, 3H, $J=7.4$ Hz); ¹³C NMR (100 MHz, CDCl₃) d 171.3, 144.8, 136.0, 132.7, 129.1, 129.0, 121.3, 78.5, 59.1, 46.2, 42.3, 32.1, 28.0, 27.8, 26.3, 21.7, 20.8, 19.8, 17.5, 13.6, 8.2; IR (neat, cm⁻¹): 3535, 2968, 2941, 2884, 1747, 1695, 1597, 1456, 1373, 1295, 1173, 1083, 814, 737; HRMS (EI) calcd for $C_{24}H_{32}N_2O_5S$ 460.2032. Found 460.2035.

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Sn(OTf)₂: β-OH/α -OH = 13:87 (62%) Zn(OTf)2: β-OH/ α-OH = 86:14 (87%)

. 17. Compounds ³ (CCDC 284467), ⁴ (CCDC 281390), 5a (CCDC 281384), 5b (CCDC 281386), 6a (CCDC 281385), 6b (CCDC 281387), 7a (CCDC 281388), 7b (CCDC 281389). TOC.

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Synthesis of Fréchet type dendritic benzyl propargyl ether and Fréchet type triazole dendrimer

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Abstract—Fréchet type dendritic benzyl propargyl ethers were synthesized by the reaction of propargyl bromide with the corresponding Fréchet type dendritic benzyl alcohol. A propargyl focal point functionalized dendrons were applied for the construction of symmetric and unsymmetric dendrimers containing 1,2,3-triazole rings as connectors via click chemistry with a tripodal azide core or a azide focal point functionalized Fréchet type dendrons.

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1. Introduction

Terminal alkynes are versatile intermediates in synthetic organic and material chemistry due to their characteristic reactions such as metathesis reaction, metal-catalyzed coupling reactions including sonogashira coupling reaction and oxidative homocoupling, and so on. Another important viewpoint in synthetic chemistry is 1,3-dipolar cycloaddition reaction with organoazides to provide heterocyclic five membered rings. Since the 1,3-dipolar cycloaddition of alkynes with azides was investigated by Huisgen et al.^{[1](#page-129-0)} it has been attracted much attention because of the synthetic importance of the aromatic and nonaromatic five-membered $[1,2,3]$ $[1,2,3]$ $[1,2,3]$ -triazole heterocycles.² The traditional method for producing the triazole by cycloaddition requires elevated temperature, typically in refluxing conditions and also provides a mixture of 1,4-disubstituted and 1,5-disubstituted triazoles.

Recently, Tornøe and Sharpless independently reported a copper(I)-catalyzed Huisgen $[2+3]$ dipolar cycloaddition reaction between an terminal alkyne and an organic azide in which the 1,4-regioisomer is exclusively formed and which also allows the rapid synthesis of compound libraries. 3 The reaction is highly chemoselective affording only the desired 1,2,3-triazole even in the presence of a large variety of other functional groups. In addition, the reaction is high yielding and can be carried out in water. The Cu(I)-catalyzed Huisgen's 1,3-dipolar cycloaddition reaction between alkynes and azides is one of the prototype reactions in click chemistry.[4](#page-129-0) This click chemistry is a modular approach that uses the most practical and reliable chemical transformations and has found in many applications in organic chemistry,^{[5](#page-129-0)} drug discovery,^{[6](#page-129-0)} bioconjugations,^{[7](#page-129-0)} material science, 8 and synthesis of polymer^{[9](#page-129-0)} and dendrimer.^{[10](#page-129-0)}

Because dendrimers contain three distinct structural parts that are the core, end-groups, and branched units connecting core and periphery, there are three strategies for triazole dendrimers. Therefore, three types of triazole dendrimer having triazole unit(s) at core(s), every branching points, and peripheries, have been synthesized convergently and/or divergently. The convergent method for dendrimer containing triazole unit(s) at core can be facilitated by fewer coupling reaction(s) between a dendron-azide and a dendron-alkyne, between a dendronazide and multi-alkynes, or between a dendron-alkyne and multi-azides and by convenient purifications. A relatively few applications using the alkynyl-dendron in dendrimer synthesis have been reported. Because of the high yields and lack of byproducts provided by the click chemistry for stitching together dendrons and core unit, the various dendrimers having functional building block at core could be obtained easily and shown the characteristic behaviors. Due to our interest in developing new functional dendrimers, we became involved in exploring efficient cycloaddition reaction that provides an easy access to dendrimers. Here we present the synthesis of propargyl-functionalized Fréchet-type

Keywords: Alkyne; Azide; Cycloaddition; Dendrimer; 1,2,3-Triazoles.

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Figure 1. Structures of acetylenic-dendrons 1-Dm $(m=1-4:$ generation of dendron).

Figure 2. Synthetic strategies of triazole dendrimers via the Cu(I)-catalyzed 1,3-dipolar cycloaddition reactions. Conditions: 5 mol% CuSO₄·5H₂O/10 mol% sodium ascorbate with respect to alkyne, DMF/H₂O (4:1), 50–60 °C.

dendrons 1-Dm (Fig. 1) and their application to the convergent synthesis of dendrimers using click chemistry with a tripodal azide core or an azide focal point functionalized Fréchet type dendrons (Fig. 2). The fundamental study reported herein details the growth of dendrimers convergently with triazole linkages between the core and dendrons.

2. Results and discussion

The poly(benzyl ether) dendrons, now frequently referred to as Fréchet-type dendrons, have been utilized by a number of groups because they are relatively readily accessed and exhibit the chemical stability associated with ether linkages and good solubility in organic solvents.^{[11](#page-129-0)} Due to these reasons we selected the Fréchet-type dendrons in the design and synthesis of the propargyl focal point functionalized dendrons. The terminal acetylenic Fréchet-type dendrons 1-Dm ($m=1-4$: generation of dendron) were synthesized by the propargylation of the corresponding dendritic benzyl alcohols with propargyl bromide [\(Scheme 1\)](#page-125-0). The Fréchettype benzyl alcohols 6-Dm $(m=1-4:$ generation of dendron) were prepared according to the reported pro-cedure.^{[12](#page-129-0)} For the synthesis of the propargyl focal point functionalized Fréchet-type dendrons 1-Dm, we have reacted the corresponding dendritic benzyl alcohols 6-Dm with propargyl bromide in the presence of NaH in THF.

Scheme 1. Synthesis of acetylenic-dendrons 1-Dm. Conditions: NaH, THF, propargyl bromide, rt, \sim 10 h.

The reaction of propargyl bromide with first generation dendritic benzyl alcohol 6-D1 in THF in the presence of NaH provided the first generation dendritic benzyl propargyl ether 1-D1 in 97% yield. Next, we conducted the reaction for the preparation of higher generation dendrons. The reactions of propargyl bromide with 6-D2, 6-D3, and 6-D4 in the same condition gave the dendritic benzyl propargyl ether 1-D2, 1-D3, and 1-D4 in yields of 92, 93, and 88%, respectively. All dendrons were confirmed by ¹H and ¹³C NMR spectroscopy, IR spectroscopy, and their FAB mass spectra.

For the construction of the triazole dendrimer 3-Gn via the 1,3-dipolar cycloaddition reactions between acetylenedendrons 1-Dm and the tripodal azide 2, we simply utilized the click chemistry condition, which is well-documented. The active Cu(I) species, generated in situ by reacting $CuSO₄·5H₂O$ with sodium ascorbate as the reducing agent, provide the 1,4-disubstituted 1,2,3-triazole in excellent yield.^{[3a](#page-129-0)} We carried out the reactions in a 4:1 solvent ratio of DMF to H_2O using 5 mol% CuSO₄ · 5H₂O with 10 mol% sodium ascorbate with respect to alkyne at $50-60$ °C. The reaction progress could be checked by TLC. The generation and disappearance of the intermediates, which are monoand/or di-triazole derivatives, were monitored by TLC runs of the reaction mixture. The reaction of 1,3,5-tris(azidomethyl)benzene 2 with 1-D1 in 0.1 M solution provided the triazole dendrimer 3-G1 having just 1,4-disubstituted 1,2,3 triazole units in yield of 89% after 18 h. Given the success in using cycloaddition reaction in the synthesis of first generation dendrimer, we expanded this reaction to get higher generation dendrimers. Reaction of 1,3,5-tris(azidomethyl)benzene 2 with 1-D2 and 1-D3 afforded the triazole dendrimers 3-G2 and 3-G3 in yields of 88 and 80%, respectively, after 26 h. In case of 1-D4, the triazole dendrimer 3-G4 was obtained in 80% yield after 28 h. For completion of the reaction between the dendritic acetylene and the tripodal core, the higher generation dendron takes longer time than the lower generation dendron. This observation led us to imagine that the reaction between the dendritic acetylene and the tripodal azide core was kinetically controlled by the accessibility of acetylide due to the steric hindrance (bulkiness) of dendron and spatial congestion of tripodal core region. Therefore, the results showed that the formation of triazole between tripodal azide and propargyl-dendrons can be regarded as a new connector to construct various dendrimers and functional materials.

All dendrimers 3-Gn were confirmed by 1 H and 13 C NMR spectroscopy. From their ${}^{1}H$ NMR spectra (CDCl₃), the peaks of the benzene protons of core and the triazole protons in dendrimers 3-Gn were found at 7.11 and 7.49 ppm for 3-G1, 7.11 and 7.49 ppm for 3-G2, 7.04 and 7.39 ppm for 3-G3, and 6.96 and 7.34 ppm for 3-G4, respectively. The peaks of the benzylic protons adjacent to the nitrogen of triazole in dendrimers 3-Gn were found at 5.43 ppm for 3-G1, 5.42 ppm for 3-G2, 5.32 ppm for 3-G3, and 5.22 ppm for 3-G4, respectively. As the dendrimer generation increased, the peaks of the benzene protons of core, the triazole protons, and the benzylic protons adjacent to the nitrogen of triazole showed up-field shift. In third and fourth generation dendrimers it is observed that the benzene protons of core, the triazole protons, and the benzylic protons adjacent to the nitrogen of triazole are influenced by the larger dendritic effect changing their microenvironment.¹³ Analysis of the dendrimers by FAB or MALDI-TOF mass spectrometry as well as by gel-permeation chromatography (GPC) provides no signs of products with defects that would arise from incomplete coupling (Fig. 3). As expected, the obtained dendrimer possessed a very well-defined molecular structure with very low polydispersity values (PDI $=1.01-1.04$). IR data also confirmed that neither alkyne $(\sim 3285 \text{ cm}^{-1})$ nor azide (2098 cm⁻¹) residues remain in the final dendrimer.

Figure 3. GPC diagrams of dendrimers 3-Gn obtained from THF eluent.

To probe the viability of our approach, we next investigated the synthesis of triazole dendrimer 5-Gmn from the coupling reactions between acetylenic-dendron 1-Dm and azido-dendron 4-Dn (entries 1–4 in [Table 1](#page-126-0)). The same reaction condition as the previous trimerization was utilized in the hetero-dimerization reaction between an azide and an alkyne. The reaction of 1-D1 with 4-D1 in a 4:1 solvent ratio of DMF to H₂O using 5 mol% CuSO₄ \cdot 5H₂O with 10 mol% sodium ascorbate at $50-60$ °C provided the just 1,4disubstituted 1,2,3-triazole product 5-G11 in 90% yield. Given the success in using cycloaddition reaction in the synthesis of first generation dendrimer, we expanded this reaction to get higher generation dendrimers (entries 2–4 in Table 1). Reactions of 1-D2 with 4-D2, 1-D3 with 4-D3, and 1-D4 with 4-D4 in a same conditions provided 1,4-disubstituted 1,2,3-triazole symmetrical dendrimers 5-G22, 5-G33, and 5-G44 in yields of 89, 94, and 95%, respectively. For completion of the reaction between two dendrons, the higher generation dendrons take slightly longer time than the lower generation dendrons which can be imagined by the simple steric hindrance of dendrons. Whereas the trimerization reaction between dendrons and core to provide 3-Gn is more sluggish than the coupling reaction between two dendrons because there are some limitation in the accessibility of acetylide due to the additional spatial congestion of core region. All symmetric dendrimers were also confirmed by H and ^{13}C NMR spectroscopy and FAB and MALDI mass spectra. From their ${}^{1}H$ NMR spectra (CDCl₃), the peaks of the benzylic protons adjacent to the nitrogen of triazole and the triazole proton in dendrimers 5-Gmn were found at 5.42 and 7.48 ppm for 5-G11, 5.42 and 7.44 ppm for 5-G22, 5.39 and 7.44 ppm for 5-G33, and 5.34 and 7.42 ppm for 5-G44, respectively. As the dendrimer generation increased, the peaks of the benzylic protons adjacent to the nitrogen of triazole and the triazole proton showed slightly up-field shift. Analysis of the higher dendrimers by gel-permeation chromatography (GPC) shows very low polydispersity values, $PDI = 1.02$ and 1.04 for 5-G33 and 5-G44, respectively (Fig. 4).

Table 1. Synthesis of triazole dendrimers 5-Gmn from azido-dendrons 1-Dm and acetylenic-dendrons 4-Dn

Entry	$1-Dm$	$4-Dn$	Rxn time (h)	Product	Yield $(\%)^a$
$\overline{1}$	$1-D1$	$4-D1$	5	5-G11	90
\overline{c}	$1-D2$	$4-D2$	6	5-G22	89
3	$1-D3$	$4-D3$	7	5-G33	94
$\overline{4}$	$1-D4$	$4-D4$	8	5-G44	95
5	$1-D1$	$4-D2$	6	5-G12	84
6	$1-D1$	$4-D3$	6	5-G13	90
7	$1-D1$	$4-D4$	7	5-G14	85
8	$1-D2$	$4-D1$	6	5-G21	85
9	$1-D2$	$4-D4$	7	5-G24	92
10	$1-D3$	$4-D1$	7	5-G31	92
11	$1-D4$	$4-D2$		5-G42	88

^a Isolated yields.

Figure 4. GPC diagrams of dendrimers 5-Gmn obtained from THF eluent.

Next, we turned our attention toward the formation of unsymmetrical 1,2,3-triazole dendrimers. We have

investigated two synthetic strategies. The first one is based on the reactions of lower generations acetylenic-dendrons 1-D1 or 1-D2 with 4-Dn (entries 5–9 in Table 1). The second strategy involves the reactions using higher generations acetylenic dendrons 1-D3 and 1-D4 (entries 10–11 in Table 1). The reactions of 1-D1 with 4-D2, 4-D3, and 4-D4 provided 1,4-disubstituted 1,2,3-triazole unsymmetrical dendrimers 5-G12, 5-G13, and 5-G14 in yields of 84, 90, and 85%, respectively. The reactions of 1-D2 with 4-D1 and 4-D4 provided 1,4-disubstituted 1,2,3-triazole unsymmetrical dendrimers 5-G21 and 5-G24 in yields of 85 and 92%, respectively. The reactions of 1-D3 with 4-D1 and of 1-D4 with **4-D2** provided 1,4-disubstituted 1,2,3-triazole unsymmetrical dendrimers 5-G31 and 5-G42 in yields of 92 and 88%, respectively. Therefore, the results showed that the formation of triazole between azide-dendrons and propargyl-dendrons are found to be an efficient connector to construct various unsymmetric dendrimers and may be applied for the synthesis of functional materials. We are currently investigated the synthesis of various unsymmetric functional dendrimers using the different kinds of dendrons. All unsymmetric dendrimers were confirmed by 1 H and 13 C NMR spectroscopy and FAB mass spectra.

3. Conclusion

We have demonstrated that the propargyl-functionalized Fréchet-type dendrons are synthesized by the propargylation of the corresponding Fréchet type dendritic benzyl alcohol and that the trimerization reactions between tripodal azide and acetylene-dendrons and the coupling reactions between azido dendrons and acetylenic dendrons lead to the formation of symmetric triazole dendrimers in high yields. Furthermore, such reactions between dendrons of different size afford unsymmetrical triazole dendrimers. This reaction may then provide an insight into designing various (un)symmetrical dendrimers such as amphiphilic dendrimers. We are currently working towards various functional dendrimers using this strategy for various applications.

4. Experimental

¹H NMR spectra were recorded on a 300 or 500 MHz NMR spectrometer using the residual proton resonance of the solvent as the internal standard. Chemical shifts are reported in parts per million (ppm). When peak multiplicities are given, the following abbreviations are used: s, singlet; d, doublet; t, triplet; q, quartet; quin, quintet; d of d, doublet of a doublet; m, multiplet; br, broad. ^{13}C NMR spectra were proton decoupled and recorded on a 75 or 125 MHz NMR spectrometer using the carbon signal of the deuterated solvent as the internal standard. EI, FAB, and MALDI mass spectra were obtained from Korea Basic Science Institute in Daegu or Daejeon and POSTECH. Flash chromatography was performed with 37–75 µm silica gel. Analytical thin layer chromatography was performed on silica plates with F-254 indicator and the visualization was accomplished by UV lamp or using an iodine chamber. Polydispersity (PDI) of dendrimers was determined by gel permeation chromatography (GPC) analysis relative to polystyrene calibration (Agilent 1100 series GPC, Plgel $5 \mu m$ MIXED-C, refractive

index detector) in THF solution. All chemicals were obtained from commercial sources and used as received, unless otherwise mentioned. THF was distilled over Na/Ph_2CO ketyl. Dendritic benzyl alcohols 6-Dm and azides $4-Dn^{12}$ $4-Dn^{12}$ $4-Dn^{12}$ and 1,3,5-tris(azidomethyl)benzene^{[14](#page-129-0)} used here were prepared according to previously reported procedure.

4.1. Synthesis of dendritic benzyl propargyl ether (1-Dm)

General procedure. Dendritic benzyl alcohol 6-Dm (1 mmol) was added to a THF (10 mL) solution of sodium hydride (1.2 mmol). After stirred under nitrogen for 30 min, propargyl bromide (1.2 mmol) was added and the mixture was stirred for \sim 10 h. The reaction mixture was poured slowly into cold brine (20 mL) and the resulting solution was extracted with EtOAc $(20 \text{ mL} \times 3)$. The combined organic phase was dried with sodium sulfate, concentrated, and purified by column chromatography (EtOAc/Hex system) to afford the desired product 1-Dn.

4.1.1. Compound 1-D1. A colorless oil; 97% yield; IR 3286, 2942, 2840, 2115, 1599, 1463, 1205, 1155, 1067 cm^{-1} ; ¹H NMR (500 MHz, CDCl₃) δ 2.47 (t, J= 2.4 Hz, 1H), 3.79 (s, 6H), 4.18 (d, $J=2.4$ Hz, 2H), 4.56 (s, 2H), 6.40 (d, $J=2.1$ Hz, 1H), 6.52 (d, $J=2.1$ Hz, 2H); ¹³C NMR (75 MHz, CDCl₃) δ 160.9, 139.6, 105.7, 100.0, 79.6, 74.6, 71.5, 57.0, 55.3; MS (EI): m/z 206 [M⁺], 166, 152, 137; HRMS (EI) Calcd for $C_{12}H_{14}O_3$: 206.0943. Found: 206.0943.

4.1.2. Compound 1-D2. A colorless oil; 92% yield; IR 3285, 2939, 2838, 2116, 1598, 1459, 1204, 1154, 1054 cm^{-1} ; ¹H NMR (500 MHz, CDCl₃) δ 2.46 (t, J= 2.4 Hz, 1H), 3.80 (s, 12H), 4.16 (d, $J = 2.4$ Hz, 2H), 4.55 (s, 2H), 4.98 (s, 4H), 6.41 (d, $J=2.1$ Hz, 2H), 6.55 (d, $J=$ 2.1 Hz, 1H), $6.57(m, 4H)$, $6.60(m, 2H)$; 13 C NMR (75 MHz, CDCl3) d 161.0, 160.0, 139.7, 139.2, 106.9, 105.2, 101.7, 100.0, 79.6, 74.7, 71.4, 70.0, 57.1, 55.4; MS (EI): m/z 478 $[M^+]$, 438, 301, 151; HRMS (EI) Calcd for $C_{28}H_{30}O_7$: 478.1992. Found: 478.1992.

4.1.3. Compound 1-D3. A colorless gum; 93% yield; IR 3284, 2938, 2838, 2117, 1598, 1458, 1203, 1153, 1050 cm^{-1} ; ¹H NMR (500 MHz, CDCl₃) δ 2.46 (t, J= 2.4 Hz, 1H), 3.79 (s, 24H), 4.16 (d, $J=2.4$ Hz, 2H), 4.55 (s, 2H), 4.97 (s, 12H), 6.41 (m, 4H), 6.53–6.60 (m, 13H), 6.67 (m, 4H); ¹³C NMR (75 MHz, CDCl₃) δ 161.0, 160.1, 160.0, 139.7, 139.2, 139.1, 106.9, 106.4, 105.2, 101.6, 100.0, 79.6, 74.7, 71.4, 70.04, 70.0, 57.1, 55.3; MS (FAB): m/z 1021.5 $[M^+]$, 966.7, 572.9 410.2, 340.2; HRMS (FAB) Calcd for $C_{60}H_{62}O_{15}$: 1022.4089. Found: 1023.4167 [M⁺ + H].

4.1.4. Compound 1-D4. A colorless gum; 88% yield; IR 3285, 2942, 2840, 2116, 1599, 1458, 1206, 1153, 1069, 1054 cm^{-1} ; ¹H NMR (500 MHz, CDCl₃) δ 2.44 (t, J= 2.4 Hz, 1H), 3.77 (s, 48H), 4.14 (d, $J=2.3$ Hz, 2H), 4.53 (s, 2H), 4.95 (s, 28H), 6.39 (m, 8H), 6.55–6.60 (m, 26H), 6.66 (m, 11H); 13 C NMR (125 MHz, CDCl₃) δ 161.0, 160.1, 160.0, 139.7, 139.2, 139.1, 107.0, 106.4, 105.2, 101.6, 100.0, 79.6, 74.7, 71.4, 70.03, 70.0, 75.1, 55.3; MS (FAB):

 m/z 2110.9 [M⁺], 1960.9, 1687.9; HRMS (FAB): Calcd for $C_{124}H_{126}O_{31}$: 2110.8283. Found: 2111.8361 [M⁺ + H].

4.2. Synthesis of 1,2,3-triazole dendrimers 3-Gn by reaction between 1,3,5-tris(azidomethyl)benzene 2 and acetylene-dendrons 1-Dm

General procedure: A solution of 1,3,5-tris(azidomethyl) benzene 2 (0.01 mmol) and acetylene-dendrons 1-Dm (0.03 mmol) in DMF–H₂O $(4.1, 1 \text{ mL})$ in the presence of 15 mol% $CuSO₄·5H₂O$ with 30 mol% sodium ascorbate was stirred at $50-60$ °C for 18–28 h. The reaction was monitored by TLC regarding on the disappearance of 1-Dm and the generation and disappearance of mono- and/or di-triazole derivatives. The reaction mixture was poured into brine (20 mL) and the resulting solution was extracted with EtOAc $(20 \text{ mL} \times 3)$. The combined organic phase was dried with sodium sulfate, concentrated, and purified by column chromatography (EtOAc/Hex system) to afford the desired product.

4.2.1. Compound 3-G1. A yellowish gum; 89% yield; IR 2923, 2854, 1597, 1465, 1203, 1154, 1067, 1051 cm⁻¹; ¹H NMR (500 MHz, CDCl₃) δ 3.75 (s, 18H), 4.51 (s, 6H), 4.62 (s, 6H), 5.43 (s, 6H), 6.36 (m, 3H), 6.48 (m, 6H), 7.11 (s, 3H), 7.49 (s, 3H); ¹³C NMR (75 MHz, CDCl₃) δ 161.3, 146.2, 140.5, 137.3, 128.0, 123.2, 106.1, 100.1, 73.0, 64.0, 55.7, 53.7; MS (FAB): m/z 862.3 [M⁺], 694.2, 647.4, 544.2; HRMS (FAB) Calcd for $C_{45}H_{51}N_9O_9$: 861.3810. Found: 862.3888 [M^+ + H]. PDI: 1.01.

4.2.2. Compound 3-G2. A yellowish gum; 88% yield; IR 2926, 2854, 1596, 1461, 1203, 1156, 1054 cm⁻¹; ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$ δ 3.77 (s, 36H), 4.50 (s, 6H), 4.61 (s, 6H), 4.94 (s, 12H), 5.42 (s, 6H), 6.39 (m, 6H), 6.52 (m, 3H), 6.55 (m, 12H), 6.57 (m, 6H), 7.11 (s, 3H), 7.44 (s, 3H); 13 C NMR (75 MHz, CDCl₃) δ 161.4, 160.4, 146.2, 140.6, 139.6, 137.3, 128.0, 123.1, 107.2, 105.7, 101.9, 100.3, 72.9, 70.4, 64.0, 55.8, 53.6; MS (FAB): m/z 1678.9 [M⁺], 663.5, 647.5; HRMS (FAM) Calcd for $C_{93}H_{99}N_9O_{21}$: 1677.6956. Found: 1678.7034 $[M^+ + H]$. PDI: 1.01.

4.2.3. Compound 3-G3. A yellowish gum; 80% yield; IR 2923, 2854, 1596, 1461, 1203, 1153, 1050 cm⁻¹; ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3)$ δ 3.75 (s, 72H), 4.48 (s, 6H), 4.59 (s, 6H), 4.92 (s, 36H), 5.32 (s, 6H), 6.38 (m, 12H), 6.49–6.64 (m, 39H), 6.64 (m, 12H), 7.04 (s, 3H), 7.39 (s, 3H); 13C NMR (125 MHz, CDCl₃) δ 161.4, 160.47, 160.37, 146.1, 140.6, 139.7, 139.5, 137.3, 131.2, 128.0, 123.1, 107.1, 106.8, 105.7, 105.4, 102.0, 101.9, 100.4, 72.9, 70.5, 70.3, 64.0, 55.8, 53.6; MS (MALDI): Calcd for $C_{189}H_{195}N_9O_{45}$: 3310.3247. Found: 3333.2024 $[M^+ + Na]$. PDI: 1.03.

4.2.4. Compound 3-G4. A yellowish solid; mp $78-80$ °C; 80% yield; IR 2933, 2839, 1599, 1462, 1203, 1157, 1051 cm^{-1} ; ¹H NMR (300 MHz, CDCl₃) δ 3.72 (s, 144H), 4.44 (s, 6H), 4.54 (s, 6H), 4.89 (s, 84H), 5.22 (s, 6H), 6.36 (m, 24H), 6.52 (m, 73H), 6.62 (m, 38H), 6.96 (s, 3H), 7.34 (s, 3H); ¹³C NMR (125 MHz, CDCl₃) δ 161.4, 160.45, 160.37, 146.2, 146.0, 140.7, 139.7, 139.63, 139.56, 137.2, 132.6, 129.0, 128.9, 127.9, 123.2, 107.1, 106.8, 105.7, 105.4, 102.0, 101.9, 101.8, 100.3, 72.8, 70.4, 70.3, 69.9, 64.0, 55.7, 55.3, 55.1, 53.5; MS (MALDI): Calcd for $C_{381}H_{387}N_9O_{93}$: 6580.15. Found: 6603.42 [M⁺ + Na]. PDI: 1.04.

4.3. Synthesis of 1,2,3-triazole dendrimers 5-Gmn by reaction between propargyl-dendrons 1-Dm and azido-dendrons 4-Dn

General procedure: A mixture of propargyl-dendrons 1-Dm (0.10 mmol) and azido-dendrons 4-Dn (0.10 mmol) in DMF–H₂O (4:1, 1 mL) in the presence of $5 \text{ mol} \%$ $CuSO₄·5H₂O$ with 10 mol% sodium ascorbate was stirred at 50 °C for \sim 8 h. The reaction was monitored by TLC regarding on the disappearance of 4-Dn. The reaction mixture was poured into brine (20 mL) and the resulting solution was extracted with EtOAc $(20 \text{ mL} \times 3)$. The combined organic phase was dried with sodium sulfate, concentrated, and purified by column chromatography (EtOAc/Hex system) to afford the desired product.

4.3.1. Compound 5-G11. A colorless oil; 90% yield; IR 2939, 2839, 1599, 1462, 1203, 1157, 1066, 1047 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ 3.75 (s, 6H), 3.78 (s, 6H), 4.52 (s, 2H), 4.65 (s, 2H), 5.42 (s, 2H), 6.37 (m, 1H), 6.40 (m, 3H), 6.49 (m, 2H), 7.48 (s, 1H); 13C NMR (75 MHz, CDCl3): d 161.4, 161.0, 145.6, 140.2, 136.7, 122.5, 106.2, 105.6, 100.5, 99.9, 72.6, 63.8, 55.5, 55.4, 54.3; MS (EI): $m/z = 399$ [M⁺], 233, 151; HRMS (EI): m/z Calcd for $C_{21}H_{25}N_3O_5$: 399.1794. Found: 399.1796.

4.3.2. Compound 5-G22. A yellowish gum; 89% yield; IR 2939, 2839, 1596, 1458, 1206, 1153, 1050 cm⁻¹; ¹H NMR (300 MHz, CDCl3): d 3.78 (s, 24H), 4.52 (s, 2H), 4.64 (s, 2H), 4.92 (s, 4H), 4.95 (s, 4H), 5.42 (s, 2H), 6.40 (m, 4H), 6.49–6.60 (m, 14H), 7.44 (s, 1H); ¹³C NMR (125 MHz, CDCl3): d 161.0, 160.96, 160.3, 159.97, 145.5, 139.2, 138.8, 136.7, 122.5, 107.2, 106.8, 105.2, 102.1, 101.5, 100.0, 99.9, 72.6, 70.1, 70.0, 63.7, 55.4, 54.1; MS (FAB): $m/z = 944.4$ $[M^+]$; HRMS (FAB): m/z Calcd for C₅₃H₅₇N₃O₁₃: 943.3891. Found: 944.3970 $[M^+ + H]$.

4.3.3. Compound 5-G33. A yellowish gum; 94% yield; IR 2939, 2840, 1597, 1456, 1203, 1154, 1051 cm⁻¹; ¹H NMR (500 MHz, CDCl3): d 3.77 (s, 48H), 4.51 (s, 2H), 4.63 (s, 2H), 4.89 (s, 4H), 4.94 (s, 20H), 5.39 (s, 2H), 6.40 (m, 6H), 6.45 (m, 2H), 6.55 (m, 24H), 6.62 (m, 4H), 6.65 (m, 6H), 7.44 (s, 1H); ¹³C NMR (75 MHz, CDCl₃): δ 161.1, 160.4, 160.2, 160.1, 160.0, 145.6, 140.4, 139.6, 139.4, 139.23, 139.2, 139.0, 136.9, 122.7, 107.2, 106.8, 106.5, 105.3, 102.2, 101.8, 101.7, 100.03, 100.0, 72.4, 70.1, 70.0, 63.7, 55.4, 54.1; MS (MALDI): Calcd for $C_{117}H_{121}N_3O_{29}$: 2031.8086. Found: 2054.7959 [M^+ + Na]. PDI: 1.02.

4.3.4. Compound 5-G44. A yellowish solid; mp $76-78$ °C; 95% yield; IR 2936, 2836, 1596, 1456, 1206, 1153, 1051 cm⁻¹; ¹H NMR (500 MHz, CDCl₃): δ 3.75 (s, 96H), 4.49 (s, 2H), 4.61 (s, 2H), 4.86 (s, 6H), 4.92 (s, 50H), 5.34 (s, 2H), 6.38 (m, 16H), 6.42 (m, 2H), 6.55 (m, 54H), 6.60 (m, 4H), 6.65 (m, 14H), 7.42 (s, 1H); 13C NMR (125 MHz, CDCl3): d 161.7, 161.4, 161.1, 160.7, 160.5, 160.4, 160.2, 145.9, 140.8, 139.8, 139.7, 139.63, 139.59, 139.3, 137.4, 123.1, 108.1, 107.5, 107.1, 106.8, 105.9, 105.7, 105.4, 102.4, 102.1, 101.9, 100.4, 72.7, 70.444, 70.37, 64.0, 55.7,

54.4; MS (MALDI): Calcd for $C_{245}H_{249}N_3O_{61}$: 4211.58. Found: $4234.70 \, [M^+ + Na]$. PDI: 1.04.

4.3.5. Compound 5-G12. A yellowish gum; 84% yield; ${}^{1}H$ NMR (300 MHz, CDCl₃): δ 3.76 (s, 6H), 3.79 (s, 12H), 4.53 (s, 2H), 4.65 (s, 2H), 4.93 (s, 4H), 5.42 (s, 2H), 6.37 (m, 1H), 6.41 (m, 2H), 6.49 (m, 4H), 6.54 (m, 4H), 6.57 (m, 1H), 7.45 (s, 1H); 13C NMR (75 MHz, CDCl3): d 161.4, 161.3, 160.8, 146.0, 140.6, 139.2, 137.1, 122.9, 107.7, 106.0, 105.6, 102.6, 100.4, 100.3, 73.0, 70.5, 64.1, 55.8, 55.7, 54.6; MS (FAB): $m/z = 672.3$ [M⁺ + H]; HRMS (FAB): m/z Calcd for $C_{37}H_{41}N_3O_9$: 671.2843. Found: 672.2921 [M⁺ + H].

4.3.6. Compound 5-G13. A yellowish gum; 90% yield; ${}^{1}H$ NMR (300 MHz, CDCl₃): δ 3.75 (s, 6H), 3.78 (s, 24H), 4.52 (s, 2H), 4.64 (s, 2H), 4.92 (s, 4H), 4.97 (s, 8H), 5.41 (s, 2H), 6.36 (m, 1H), 6.41 (m, 4H), 6.46–6.49 (m, 4H), 6.57 (m, 11H), 6.54 (m, 4H), 7.46 (s, 1H); 13C NMR (75 MHz, CDCl3): d 161.7, 161.4, 161.3, 160.7, 160.5, 160.49, 160.4, 146.0, 140.7, 139.7, 139.6, 139.5, 139.3, 137.1, 122.9, 107.7, 107.2, 106.83, 106.8, 106.5, 106.0, 105.7, 102.1, 102.0, 101.9, 100.8, 100.38, 100.35, 100.27, 72.9, 72.8, 70.5, 70.4, 64.1, 55.8, 55.7, 54.6; MS (FAB): $m/z = 1216.4$ $[M^+ + H]$, 663.5; HRMS (FAB): m/z Calcd for $C_{69}H_{73}N_3O_{17}$: 1215.4940. Found: 1216.5018 [M⁺ + H].

4.3.7. Compound 5-G14. A yellowish gum; 85% yield; ${}^{1}H$ NMR (300 MHz, CDCl₃): δ 3.72 (s, 6H), 3.77 (s, 48H), 4.50 (s, 2H), 4.63 (s, 2H), 4.91 (s, 4H), 4.96 (s, 24H), 5.38 (s, 2H), 6.35 (m, 1H), 6.40 (m, 8H), 6.48 (m, 3H), 6.56 (m, 22H), 6.63 (m, 4H), 6.67 (m, 10H), 7.45 (s, 1H); ¹³C NMR $(125 \text{ MHz}, \text{CDCl}_3): \delta$ 161.0, 160.9, 160.3, 160.1, 145.6, 140.2, 139.2, 139.1, 138.7, 136.8, 122.6, 107.3, 106.4, 105.6, 105.2, 102.0, 101.7, 101.6, 99.9, 99.8, 99.6, 72.5, 70.04, 70.0, 63.6, 55.33, 55.27, 54.1; MS (FAB): $m/z =$ 1216.4 $[M^+ + H]$, 663.5; MS (FAB): Calcd for $C_{133}H_{137}N_3O_{33}$: 2305.5. Found: 2305.9 [M⁺].

4.3.8. Compound 5-G21. A yellowish gum; 85% yield; 1 H NMR (300 MHz, CDCl₃): δ 3.73 (s, 6H), 3.78 (s, 12H), 4.51 (s, 2H), 4.63 (s, 2H), 4.95 (s, 4H), 5.41 (s, 2H), 6.40 (m, 5H), 6.53–6.59 (m, 7H), 7.47 (s, 1H); ¹³C NMR (125 MHz, CDCl3): d 161.3, 161.0, 160.7, 160.0, 145.5, 140.3, 139.2, 139.1, 136.7, 136.5, 122.5, 106.7, 106.1, 105.9, 105.2, 105.0, 101.5, 100.4, 99.9, 72.4, 70.0, 63.7, 55.4, 55.3, 54.2; MS (FAB): $m/z=672.3$ [M⁺+H]; HRMS (FAB): m/z Calcd for $C_{37}H_{41}N_3O_9$: 671.2843. Found: 672.2921 [M^+ + H].

4.3.9. Compound 5-G24. A yellowish gum; 92% yield; ${}^{1}H$ NMR (300 MHz, CDCl₃): δ 3.75 (s, 12H), 3.78 (s, 48H), 4.49 (s, 2H), 4.62 (s, 2H), 4.91 (s, 8H), 4.95 (s, 24H), 5.39 (s, 2H), 6.40 (m, 10H), 6.47 (m, 2H), 6.56 (m, 30H), 6.62 (m, 4H), 6.66 (m, 8H), 7.44 (s, 1H); 13C NMR (75 MHz, CDCl3): d 161.4, 161.36, 160.7, 160.5, 160.4, 146.0, 140.7, 139.7, 139.6, 139.55, 139.3, 137.3, 123.0, 107.6, 107.1, 106.8, 106.1, 105.7, 102.5, 102.0, 101.9, 100.3, 72.8, 70.5, 70.4, 64.0, 55.7, 54.5; MS (FAB): Calcd for $C_{149}H_{153}N_3O_{37}$: 2577.8. Found: 2577.4 [M⁺].

4.3.10. Compound 5-G31. A yellowish gum; 92% yield; 1 H NMR (500 MHz, CDCl₃): δ 3.73 (s, 6H), 3.78 (s, 24H), 4.52 (s, 2H), 4.64 (s, 2H), 4.96 (s, 4H), 4.98 (s, 8H), 5.40 (s, 2H),

6.39–6.41 (m, 7H), 6.52 (m, 1H), 6.57 (m, 13H), 6.67 (m, 3H), 7.46 (s, 1H); ¹³C NMR (125 MHz, CDCl₃): δ 161.7, 161.4, 160.5, 160.4, 145.9, 140.7, 139.7, 139.6, 137.1, 122.9, 107.2, 106.8, 106.6, 105.7, 102.0, 101.9, 100.9, 100.4, 72.9, 70.5, 70.4, 64.1, 55.8, 54.6; MS (FAB): $m/z =$ 1216.4 $[M^+ + H]$, 753.5, 647.5; HRMS (FAB): m/z Calcd for $C_{69}H_{73}N_3O_{17}$: 1215.4940. Found: 1216.5018 [M⁺ + H].

4.3.11. Compound 5-G42. A yellowish gum; 88% yield; 1 H NMR (300 MHz, CDCl₃): δ 3.75 (s, 12H), 3.77 (s, 48H), 4.52 (s, 2H), 4.63 (s, 2H), 4.87 (s, 8H), 4.95 (s, 24H), 5.36 (s, 2H), 6.40 (m, 10H), 6.45 (m, 2H), 6.51 (m, 4H), 6.56 (m, 24H), 6.60 (m, 2H), 6.66 (m, 12H), 7.43 (s, 1H); ¹³C NMR (75 MHz, CDCl3): d 161.1, 160.6, 160.4, 160.1, 160.0, 145.6, 140.4, 139.4, 139.3, 139.2, 138.9, 136.9, 122.6, 107.6, 107.2, 106.8, 106.5, 105.3, 102.2, 102.0, 101.7, 100.0, 72.5, 70.1, 69.8, 63.7, 55.4, 54.1; MS (FAB): Calcd for $C_{149}H_{153}N_3O_{37}$: 2577.8. Found: 2577.4 [M⁺].

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Synthesis of atropoisomeric pyridines via cobalt-catalyzed cocyclotrimerization of diynes with benzonitrile

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Abstract—Arylpyridines (precursors for potential organocatalysts) are easily accesible by cobalt-catalyzed cocyclotrimerization of *ortho*substituted 1-aryl-1,7-octadiynes with benzonitrile. The scope of the reaction with respect to the ortho substituents (OMe, Me, COOMe, NHCOMe, F, etc.) was investigated. Three potentially atropoisomeric arylpyridines were prepared and one of them was converted into the corresponding N-oxide and resolved into its enantiomers. The absolute configuration of the N-oxide was established by X-ray crystal structure analysis. Preliminary results of its application in asymmetric organocatalysis are presented. Q 2005 Elsevier Ltd. All rights reserved.

1. Introduction

The interest in the field of organocatalysis (acceleration of a reaction by a catalytic amount of an organic compound) has increased in the last few years. In particular, organocatalysis is gaining importance in asymmetric synthesis, comple-menting bio- and metal-catalysis.^{[1](#page-137-0)} Out of several concepts of organocatalysis, a significant role is played by activation of a Lewis acid by a Lewis base. One type of the typical Lewis base organocatalysts are pyridine N-oxides with a biaryl framework.^{[2–5](#page-137-0)} Since pyridine N-oxides are easily accessible from pyridines, there is general interest in the development of new synthetic methods for pyridine preparation. One of them is based on $[2+2+2]$ cocyclotrimerization of two $C-C$ -triple bonds with a nitrile.^{[6](#page-137-0)} The use of the most widely and generally utilized cobalt

catalysts was pioneered by Wakatsuki,⁷ Vollhardt,^{[8](#page-137-0)} and Bönnemann. 9 Over the years a number of other transition metal compounds such as $Ti¹⁰, Zr¹¹, Fe¹², Ta¹³$ and Rh^{[14](#page-137-0)} were shown to catalyze or mediate the cyclotrimerization. Recently, a Ru-based catalyst has been shown to be suitable for cyclotrimerization of diynes with electron-deficient nitriles.[15,16](#page-137-0) Cocyclotrimerization has also been used for preparation of oligopyridines (namely bipyridines) either by cocyclotrimerization of diynes with dinitriles, 17 or alkynyl-nitriles with diynes,^{[18](#page-137-0)} or cyanopyridines with alkynes.^{[19](#page-137-0)} The synthesis of chiral pyridines is based either on cyclotrimerization with chiral nitriles^{[20](#page-137-0)} or enantioselective cyclotrimerization by treatment with chiral cyclopenta d dienyl cobalt complexes.^{[21](#page-137-0)} Herein, we report on the cobaltcatalyzed cyclotrimerization of substituted aryldiynes with nitriles to potentially atropoisomeric pyridines, conversion

Scheme 1. Preparation of diynes 2 and their cocyclotrimerization with nitriles to arylpyridines 3 under Co-catalysis.

Keywords: Pyridine; Cocyclotrimerization; Cobalt; Catalysis; Organocatalysis.

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^a Conditions: A = CpCo(CO)₂ (20 mol%), PPh₃ (40 mol%), 140 °C, 48 h; B = CpCo(COD) (20 mol%), 140 °C, 48 h; C = CpCo(CH₂ = CH₂)₂ (10 mol%), $20 °C$, 30 min.
^b Isolated yields.

 $c^{\rm c}$ CpCo(CO)₂ (10 mol%), 140 °C, 48 h.

of one of them into a pyridine N-oxide, its resolution into enantiomers, and preliminary results of its application in enantioselective additions to benzaldehyde.

2. Results and discussion

We envisioned that one of the possible pathways to potentially atropoisomeric arylpyridines could be based on the reaction of nitriles with properly substituted α , ω -diynes,^{[22,23](#page-137-0)} such as 1-(ortho-substituted-aryl)-1,7-octadiynes [\(Scheme 1](#page-130-0)). This strategy is similar to that used by Heller et al.^{[21](#page-137-0)} but its scope is limited only to a methoxy group as the substituent in the ortho-position. It is reasonable to assume that ortho-substitution may affect the course of cyclotrimerization by steric and electronic effects. These effects are important in the search for new synthetic methods for preparation of potentially atropoisomeric biaryls. Recently, they have been studied in the Dötz reaction of ortho-substituted arylalkynes with chromium carbenes^{[24](#page-138-0)} and in the CuCl mediated reaction of zirconacyclopentadienes with *ortho*-substituted arylpropynoates.^{[25](#page-138-0)} Furthermore, the synthetic usefulness of the commercially available $CpCo(CO)_2$ was explored.

The required ortho-substituted 1-aryl-1,7-octadiynes were prepared by palladium-catalyzed Krause modification^{[26](#page-138-0)} of the Sonogashira coupling^{[27](#page-138-0)} of 1,7-octadiyne with arylhalides 1 ([Scheme 1\)](#page-130-0). Usually, the coupling proceeded smoothly to afford the desired diynes 2 in moderate to reasonable isolated yields (24–62%) and selectivity. To accomplish the cyclotrimerization of the diynes with nitriles, the standard conditions (20 mol% of commercially available $CpCo(CO)_2$, 40 mol% of PPh₃ as a ligand, 140 °C, 2 days) were used ([Scheme 1\)](#page-130-0). It has been shown that the catalytically active species are formed in a reasonable reaction rate at this temperature.^{[6a](#page-137-0)} Other cobalt complexes such as $CpCo(COD)$ and $CpCo(CH_2=CH_2)_2$ (Jonas catalyst)^{[28,29](#page-138-0)} were used in order to compare the catalytic activity. The cocyclotrimerizations were carried out in benzonitrile as solvent to minimize homocyclotrimerization of diyne, to ensure a high selectivity ratio for pyridine formation.

The results of cocyclotrimerizations are presented in [Table 1](#page-131-0). The reaction with the diyne bearing an ester group 2a afforded the corresponding product 3a in acceptable yield of 54% (entry 1). In the case of the diyne with a nitrogen substituent such as 2b, the reaction proceeded only in moderate yield of 35% (entry 2). The cocyclotrimerizations with alkynes bearing methoxy 2c and methoxymethyl substituents 2d differed considerably (entries 3 and 4); the product 3c was obtained in good yield of 76%, whereas the arylpyridine 3d was isolated in significantly lower yield of 33%. In the case of 1-(orthotolyl)-1,7-octadiyne 2e the arylpyridine 3e was furnished in high yield of 91% (entry 5). The cocyclotrimerization of the fluorine substituted diyne 2f with benzonitrile also proceeded to give the corresponding arylpyridine 3f in moderate 46% yield (entry 6).

Cocyclotrimerizations of diynes that were expected to give sterically hindered arylpyridines with restricted rotation about the bond connecting two aromatic rings were far more intriguing (entries 7–9). The reaction of the methoxymethyl substituted diyne 2g afforded the desired product 3g in 30% yield (entry 7, conditions A). The use of CpCo(COD) under the same reaction conditions afforded the product in a similar yield of 25% (conditions B). A considerably different result was obtained when the Jonas catalyst (condition C) was used. The full conversion of the starting material was observed within 30 min and 3g was isolated in good 62% yield. The structure of 3g was unequivocally confirmed by crystalographic analysis (Fig. 1).

Figure 1. An ORTEP diagram of 3g. Displacement parameters are shown at the 50% probability level.

The cyclotrimerization of diyne 2g was also carried out on 10 mmol scale and the product 3g was isolated in 45% yield. In addition, it was possible to isolate the compound 4 in 6% yield, which was the product of the cocyclotrimerization of two molecules of 2g with benzonitrile ([Scheme 2\)](#page-133-0). As expected, the reaction with the methoxynaphthyldiyne 2h furnished the corresponding product 3h in very good yield of 75% (entry 8, conditions A), which is close to the yield obtained by Heller under different reaction conditions.^{[21](#page-137-0)} The similar result was observed under conditions B (67% yield). Surprisingly, the reaction carried out in the presence of the Jonas catalyst did not proceed and the starting material was quantitatively recovered (entry 8, conditions C). The reaction with methylnaphthyldiyne 2i gave the corresponding product 3i in rather low yield of 30% (entry 9, conditions A). Again the use the Jonas catalyst (conditions C) resulted in increase of the reaction rate and the conversion of the starting material; the product 3i was isolated in 48% yield. However, in this instance CpCo(COD) proved to be the catalyst of choice, because

Scheme 2. Formation of 4 in cocyclotrimerization of 2g and benzonitrile.

its use afforded the corresponding product in 69% yield (conditions B).

The question is what factors are mainly responsible for the observed difference in the yields of the arylpyridines in entries 1–6. Obviously, the differences cannot be simply explained by steric factors, since it is known that for

Scheme 3. Oxidation of the pyridine 3g to 6.

example methyl group occupies larger space than methoxy or ester groups.[30](#page-138-0) Although we do not have any spectroscopic evidence for the following hypothesis, it is sensible to assume that the course of the reaction is influenced (retarded) by the strengh of the coordination of the lone electron pair on the heteroatom of the ortho-substituent to the cobalt atom in the intermediate cobaltacyclopentadiene.

Since our initial impetus for this work was to develop an alternative method for the preparation of chiral pyridine oxides as potential organocatalysts, we chose pyridine 3g for further investigation in this direction. Its oxidation with m-chloroperoxobenzoic acid (MPCBA) proceeded smoothly to give the corresponding N-oxide 6 in 54% isolated yield (Scheme 3). The N-oxide 6 was resolved by cocrystallization with (S) - $(-)$ -binol 7 (binol=2,2'-di-hydroxy-1,1'-binaphthyl),^{[2–5](#page-137-0)} which gave the crystalline

Figure 2. An ORTEP diagram illustrating the interaction of (R) -(+)-6 (on the left) with (S) -(-)-7 (right), in particular the hydrogen bonding N–O···H–O; $O(1)\cdots O(4a)$ 2.606(2) \AA , $O(1)\cdots H-O(4a)$ 176(2)°. Displacement parameters are shown at the 50% probability level.

material containing $(S)-(-)$ -binol 7 and $(+)$ -6 (in 1:1) ratio), while $(-)$ -6 remained in the solution. This cocrystallization, followed by a chromatographic separation of $(+)$ -6 from (S) - $(-)$ -binol 7, furnished pure $(+)$ -6 of 95% ee (as detected by chiral HPLC, Chiracel OD-H) in 30% yield. The absolute configuration was found to be (R) - $(+)$ -6 by crystallographic analysis of the molecular crystal of $(+)$ -6 with (S) - $(-)$ -7 ([Fig. 2](#page-133-0)) of known absolute configuration. The configurational stability of the compound $(R)-(+)$ -6 was quantitatively evaluated by an analytical chiral HPLC. Preliminary data were obtained by heating samples of $(R)-(+)$ -6 and injecting them on the chiral column. The $(R)-(+)$ -6 showed no racemization in toluene at 110° C after 12 h.

The catalytic activity of the pyridine N-oxide $(R)-(+)$ -6 (5 mol%) was preliminarly tested in the addition of allytrichlorosilane 8 to benzaldehyde 7 in dichloromethane (Scheme 4). The attempt to carry out the reaction under usual conditions $(-40 \degree C)^5$ $(-40 \degree C)^5$ was not successful. The reaction proceeded at room temperature only, but even then the reaction rate was rather low, 50% yield of the corresponding alcohol 9 was obtained after 72 h with the modest asymmetric induction of 20% ee. The enantioselectivity of $(R)-(+)$ -6 was also tested in the reaction of diethylzinc $\overline{10}$ with benzaldehyde $\overline{7}^{31}$ $\overline{7}^{31}$ $\overline{7}^{31}$ The resulting asymmetric induction was again modest (17% ee).

Scheme 4. Enantioselective additions to benzaldehyde 7.

3. Conclusion

This work has shown that cobalt-catalyzed $[2+2+2]$ cocyclotrimerization of diynes with benzonitrile is a convenient and straightforward method for preparation of not only mono-ortho-substituted arylpyridines but also potentially atropoisomeric bis-ortho-substituted arylpyridines (entries 7–9, [Table 1\)](#page-131-0) in reasonable yields. The starting diynes are easily prepared from the corresponding substituted arylhalides and 1,7-octadiyne in good yields and the use of the commercially available $CpCo(CO)_2$ catalysts gave good yields of the cyclotrimerization products in most cases. Other types of $CpCo(ligand)_n$ (ligand=ethylene, COD) catalysts were tested in cyclotrimerization with bis-orthosubstituted aryldiynes but their activity was highly dependent on the structure of the substrate. Last but not least, the racemic arylpyridine-N-oxide 6 was resolved into enantiomers and its absolute configuration was unequivocally determined by X-ray structure analysis. Despite the fact that the result of enantioselective additions were not too high, it is premature to rule out the synthetic utility of the pyridine N-oxide $(R)-(+)$ -6 at this moment. Further experiments investigating its reactivity and structural modifications will follow in the near future.

4. Experimental

All reactions were carried out under a protective atmosphere of Ar in 20 mL Schlenk flasks. Unless mentioned, reagents were used as obtained without further purification.

4.1. General procedure for catalytic cyclotrimerization of aryl-substituted 1,7-octadiynes with benzonitrile

1-Aryl-1,7-octadiyne (0.4 mmol) and PPh₃ (42 mg) , 0.16 mmol) were added to a dry Schlenk tube under argon and dissolved in benzonitrile (2 mL, 19.4 mmol). Then $CpCo(CO)$ ₂ (14 mg, 0.08 mmol) was added. The reaction mixture was heated for 48 h at 140° C. After cooling down it was quenched with water and volatiles were removed under reduced pressure. Column chromatography on silica gel was used to isolate products.

4.1.1. Methyl 2-(5,6,7,8-tetrahydro-3-phenylisoquinolin-1-yl)benzoate (3a). Column chromatography on silica gel (7:1 hexane/EtOAc) afforded 192 mg (54%) of the title compound as a viscous liquid: ${}^{1}H NMR$ (400 MHz, CDCl₃) δ 1.74–1.84 (m, 4H), 2.44–2.47 (m, 2H), 2.87–2.90 (m, 2H), 3.61 (s, 3H), 7.30–7.47 (m, 6H), 7.54–7.59 (m, 1H), 7.93–7.95 (m, 2H), 7.99–8.01 (m, 1H); ¹³C NMR (100 MHz, CDCl3) d 22.4, 23.0, 26.7, 29.5, 52.0, 119.8, 126.9 (2C), 127.7, 128.1, 128.4 (2C), 129.4, 130.0, 130.1, 130.3, 131.6, 139.7, 142.3, 146.6, 153.1, 159.0, 167.7; IR (CHCl₃) ν 3523, 3367, 3064, 3005, 2952, 1721, 1590, 1576, 1434, 1295, 1275, 1130, 1084, 1052, 966, 908 cm⁻¹; FAB-MS m/z 241 (M+H⁺), 225, 165, 128, 115; HR-MS (FAB) calculated for $C_{16}H_{17}O_2$ (M+H⁺) 241.1239, found 241.1229.

4.1.2. N-(2-(5,6,7,8-Tetrahydro-3-phenylisoquinolin-1 yl)phenyl)acetamide (3b). Column chromatography on silica gel (1:1 hexane/EtOAc) afforded 96 mg (35%) of the title compound as a pale yellow solid: mp $130-132$ °C (hexane); ¹H NMR (400 MHz, CDCl₃) δ 1.72–1.76 (m, 2H), 1.83–1.89 (m, 2H), 1.92 (s, 3H), 2.66–2.69 (m, 2H), 2.92–2.95 (m, 2H), 7.13–7.17 (m, 1H), 7.35–7.51 (m, 6H), 8.01–8.03 (m, 2H), 8.33 (d, $J=8.4$ Hz, 1H), 9.61 (br s, 1H); ¹³C NMR (100 MHz, CDCl₃) δ 22.1, 22.9, 24.8, 27.6, 29.7, 119.7, 122.4, 123.0, 126.2 (2C), 128.4, 128.7 (2C), 128.8, 128.9, 129.9, 131.5, 135.8, 138.5, 149.5, 152.5, 156.0, 168.1; IR (CHCl₃) ν 3325, 3065, 3011, 2943, 2864, 1682, 1590, 1521, 1448, 1303, 1242 cm⁻¹; EI-MS m/z (% relative intensity) 342 (M^+ , 100), 327 (95), 299 (28), 284 (20), 271 (9), 225 (16), 171 (8), 149 (32), 105 (86), 77 (52), 57 (26), 55 (19), 43 (27); HR-MS calculated for $C_{23}H_{22}N_{2}O$ 342.1732, found 342.1749.

4.1.3. 5,6,7,8-Tetrahydro-1-(2-methoxyphenyl)-3-phenylisoquinoline (3c). Column chromatography on silica gel $(4:1)$ hexane/EtOAc) afforded 97 mg $(76%)$ of the title compound as a viscous liquid: ${}^{1}H NMR$ (400 MHz, CDCl₃) δ 1.65–1.85 (m, 4H), 2.33–2.40 (m, 1H), 2.60–2.66 (m, 1H), 2.85–2.90 (m, 2H), 3.77 (s, 3H), 6.95–6.97 (m, 1H), 7.04–7.07 (m, 1H), 7.30–7.42 (m, 6H), 7.96–7.98 (m, 2H); ¹³C NMR (100 MHz, CDCl₃) δ 22.4, 22.9, 25.7, 29.6, 55.5, 110.8, 120.0, 120.7, 127.0 (2C), 128.1, 128.4 (2C), 129.2, 130.4, 130.7, 131.0, 139.9, 146.5, 153.6, 156.6, 157.0; IR (CHCl₃) ν 3530, 3063, 2940, 2863, 2837, 1697, 1589, 1555, 1495, 1463, 1278, 1244, 1180, 1109, 1027, 955 cm⁻¹; EI-MS m/z (% relative intensity) 315 (M⁺, 100), 298 (10), 284 (30), 210 (13); HR-MS calculated for $C_{22}H_{21}NO$ 315.1623, found 315.1638.

4.1.4. 5,6,7,8-Tetrahydro-1-(2-(methoxymethyl)phenyl)- 3-phenylisoquinoline (3d). Column chromatography on silica gel (5:2 hexane/EtOAc) afforded 56 mg (33%) of the title compound as a viscous liquid: ${}^{1}H$ NMR (400 MHz, CDCl3) d 1.71–1.88 (m, 4H), 2.35–2.58 (m, 2H), 2.87–2.91 (m, 2H), 3.25 (s, 3H), 4.33 (s, 3H), 7.16–7.43 (m, 7H), $7.56-7.57$ (m, 1H), $7.95-7.97$ (m, 2H); 13 C NMR $(100 \text{ MHz}, \text{ CDCl}_3)$ δ 22.3, 23.0, 26.7, 29.6, 58.3, 72.2, 119.9, 125.3, 126.8, 127.2, 127.9, 128.1, 128.2, 128.3, 128.5 (2C), 128.8, 129.0, 130.2, 136.3, 139.4, 153.1, 158.3; IR $(CHCl₃)$ v 3063, 3008, 2935, 2862, 1589, 1432, 1090, 956 cm⁻¹; EI-MS m/z (% relative intensity) 329 (M⁺, 16), 314 (100), 277 (9), 149 (11), 69 (25), 55 (20), 43 (23); HR-MS calculated for $C_{23}H_{23}NO$ 329.1780, found 329.1768.

4.1.5. 5,6,7,8-Tetrahydro-3-phenyl-1-o-tolylisoquinoline (3e). Column chromatography on silica gel (4:1 hexane/ EtOAc) afforded 110 mg (91%) of the title compound as a viscous liquid: ¹H NMR (400 MHz, CDCl₃) δ 1.71–1.83 (m, 4H), 2.14 (s, 3H), 2.30–2.45 (m, 2H), 2.86–2.89 (m, 2H), 7.19–7.29 (m, 4H), 7.33–7.42 (m, 4H), 7.95–7.98 (m, 2H); ¹³C NMR (100 MHz, CDCl₃) δ 19.6, 22.3, 23.0, 26.5, 29.6, 119.7, 125.5, 126.8 (2C), 127.6, 128.2, 128.5 (2C), 128.6, 129.7, 130.2, 135.7, 139.8, 140.5, 147.1, 153.5, 159.5; IR (CHCl₃) ν 3523, 3063, 2942, 2863, 1692, 1589, 1580, 1554, 1432, 1387, 1247, 1178, 1072, 1026 cm⁻ ; EI-MS m/z (% relative intensity) 299 (M⁺, 100), 284 (65), 270 (20), 257 (26), 165 (9), 128 (8), 103 (8), 84 (27), 49 (7); HR-MS calculated for $C_{22}H_{21}N$ 299.1674, found 299.1672.

4.1.6. 5,6,7,8-Tetrahydro-1-(2-fluorophenyl)-3-phenylisoquinoline (3f). Column chromatography on silica gel (4:1 hexane/EtOAc) afforded 122 mg (46%) of the title compound as a viscous liquid: ¹H NMR (400 MHz, C_6D_6) δ 1.41–1.46 (m, 4H), 2.41–2.50 (m, 4H), 2.60–2.66 (m, 1H), 6.63–6.67 (m, 1H), 6.79–6.83 (m, 1H), 6.90–6.96 (m, 2H), 7.16–7.20 (m, 1H), 7.24–7.29 (m, 2H), 7.40–7.45 (m, 1H), 8.16–8.19 (m, 2H); ¹³C NMR (100 MHz, C₆D₆) δ 23.3, 23.8, 27.0, 30.3, 116.9, 121.5, 125.7 (d, $J=3$ Hz), 128.6 $(2C)$, 130.2 $(2C)$, 131.2, 132.4, 133.4 $(d, J=18$ Hz), 141.4, 148.5, 155.5, 156.2, 161.7 (d, $J=246$ Hz); IR (CHCl₃) ν 2941, 1713, 1617, 1590, 1580, 1553, 1494, 1452, 1432, 1423, 1386, 1226, 1099 cm⁻¹; EI-MS m/z (% relative intensity) 303 (35), 133 (10), 103 (M^+ , 100), 76 (32), 50 (16); HR-MS calculated for $C_{21}H_{18}FN$ 303.1423, found 303.1435.

4.1.7. 5,6,7,8-Tetrahydro-1-(2-methoxy-6-methylphenyl)-3-phenylisoquinoline (3g). Column chromatography on silica gel (4:1 hexane/EtOAc) afforded 121 mg (30%) of the title compound as a white solid: mp

138–139 °C (hexane); ¹H NMR (400 MHz, CDCl₃) δ 1.69–1.83 (m, 4H), 2.03 (s, 3H), 2.04–2.27 (m, 1H), 2.43– 2.49 (m, 1H), 2.86–2.89 (m, 2H), 3.71 (s, 3H), 6.81 (d, $J=$ 8.4 Hz, 1H), 6.91 (d, $J=7.6$ Hz, 1H), 7.23–7.27 (m, 1H), 7.31–7.42 (m, 4H), 7.94–7.97 (m, 2H); ¹³C NMR (100 MHz, CDCl3) d 19.7, 22.3, 22.8, 25.4, 29.6, 55.6, 108.4, 120.1, 122.7 (2C), 127.1, 128.2, 128.3, 128.4 (2C), 128.6, 131.1, 137.6, 139.8, 146.9, 153.7, 156.4, 156.7; IR (CHCl₃) v 3065, 3025, 3016, 2941, 2863, 2839, 2357, 1591, 1581, 1556, 1470, 1434, 1422, 1387, 1297, 1261, 1230, 1221, 1212, 1087 cm⁻¹; EI-MS m/z (% relative intensity) 329 (MC, 29), 314 (20), 298 (12), 284 (10), 256 (15), 242 (6), 148 (31), 135 (9), 123 (10), 111 (13), 97 (24), 81 (42), 69 (100), 57 (67), 43 (77); HR-MS calculated for $C_{23}H_{23}NO$ 329.1780, found 329.1784. EA calculated for $C_{23}H_{23}NO$ C, 83.85; H, 7.04; N, 4.25. Found C, 83.59; H, 7.08; N, 3.99.

4.1.8. 5,6,7,8-Tetrahydro-1-(2-methoxynaphthalen-1-yl)- 3-phenylisoquinoline (3h). Spectral characteritics were in agreement with the previously published data. 21 21 21

4.1.9. 5,6,7,8-Tetrahydro-1-(2-methylnaphthalen-1-yl)- 3-phenylisoquinoline (3i). Column chromatography on silica gel (10:1 hexane/EtOAc) afforded 125 mg (30%) of the title compound as a white solid: mp $175-176$ °C (hexane); ¹H NMR (400 MHz, CDCl₃) δ 1.63–1.69 (m, 2H), 1.77–1.83 (m, 2H), 2.13–2.22 (m, 2H), 2.22 (s, 3H), 2.91–2.94 (m, 2H), 7.22–7.24 (m, 1H), 7.29–7.50 (m, 7H), 7.79–7.84 (m, 2H), 7.96–7.98 (m, 2H); ¹³C NMR $(100 \text{ MHz}, \text{CDCl}_3)$ δ 20.0, 22.3, 22.9, 25.6, 29.7, 120.1, 124.7, 125.1, 126.0, 126.9 (2C), 127.6, 127.9, 128.3, 128.5 (2C), 128.8, 131.0, 131.9, 132.1, 133.1, 136.6, 139.8, 147.2, 154.3, 158.1; IR (CHCl₃) ν 3059, 3024, 3015, 2941, 2356, 1591, 1556, 1508, 1494, 1381, 1315, 1260, 1229, 1219, 1205, 1198, 1085, 915, 866, 813 cm⁻¹; EI-MS m/z (% relative intensity) 349 (M^+ , 100), 334 (65), 320 (16), 307 (14), 256 (7), 166 (8), 149 (46), 139 (8), 111 (10), 97 (16), 83 (24), 69 (42), 57 (56), 43 (59); HR-MS calculated for $C_{26}H_{23}N$ 349.1830, found 349.1839. EA calculated for $C_{26}H_{23}N$ C, 89.36; H, 6.63; H 4.01. Found C, 88.86; H, 6.67; H, 3.76.

4.1.10. $(+)$ -5,6,7,8-Tetrahydro-1-(2-methoxy-6-methylphenyl)-3-phenylisoquinoline-N-oxide (6). To a solution of pyridine 3f (160 mg, 0.48 mmol) in dichloromethane (2 mL) was added MCPBA (purity 70%) (220 mg, 0.93 mmol) at 0° C. After stirring of the resulting mixture at room temperature for 1 h it was quenched by the saturated water solution of NaHCO₃ (1 mL) and the crude product was extracted with dichloromethane (5 mL). The organic layer was separated and column chromatography on silica gel (ethyl acetate) afforded 90 mg (54%) of the title compound 6 as a viscous liquid: H NMR (400 MHz, C_6D_6) δ 1.32–1.44 (m, 4H), 2.06–2.14 (m, 2H), 2.16 (s, 3H), 2.26–2.33 (m, 2H), 3.25 (s, 3H), 6.56 (d, $J=8.2$ Hz, 1H), 6.91 (d, $J=7.6$ Hz, 1H), $7.01-7.06$ (m, 1H), $7.10-7.22$ (m, 4H), 7.96–7.98 (m, 2H); ¹³C NMR (100 MHz, CDCl₃) δ 19.9, 23.1, 23.3, 27.1, 29.2, 56.2, 109.8, 123.6, 123.9, 126.8, 128.6 (2C), 129.5, 130.3, 130.6 (2C), 134.0, 134.7, 135.4, 139.5, 146.9, 148.0, 158.0; IR (CHCl₃) ν 3614, 3010, 2971, 1582, 1467, 1387, 1257, 1127, 1080, 945 cm⁻¹; EIS-MS m/z 346 (M⁺ +H), 352 (M⁺ -O+Na), 368 (M⁺ +Na).

4.1.11. $(R)-(+)$ -5,6,7,8-Tetrahydro-1-(2-methoxy-6methylphenyl)-3-phenylisoquinoline-N-oxide $((R)-(+)$ -6). To a solution of $(S)-(-)$ -binol 7 (293 mg, 0.99 mmol) and racemic 5,6,7,8-tetrahydro-1-(2-methoxy-6-methylphenyl)- 3-phenylisoquinoline-N-oxide 6 (340 mg, 0.99 mmol) in dichloromethane (5 mL) heptane (10 mL) was added, the flask was closed with a septum with a needle and set aside to allow slow evaporation of dichloromethane through the needle. The molecular complex $(R)-(+)$ -6 \cdot (S)-(-)-7 crystallized within 5 days as colorless needles that were collected by suction filtration. Individual components were collected by column chromatography on silica gel (ethyl acetate), which afforded 105 mg (30%) of $(R)-(+)$ -6. Chiral HPLC (Chiralcel OD-H, 0.46×25 cm, 8:1 heptane/ 2-propanol, 1.2 mL min⁻¹) showed 95% ee depending on the batch ($t_S = 8.01$ min, $t_R = 9.12$ min).

4.2. Enantioselective allylation of benzaldehyde with allyltrichlorosilane to $(R)-(+)$ -1-phenyl-but-3-en-1-ol (9)

To a solution of $(R)-(+)$ -6 (5 mg, 0.014 mmol) in dichloromethane (1.4 mL) were added benzaldehyde $(40 \mu L,$ 0.4 mmol), diisopropylethylamine $(87 \mu L, 0.5 \text{ mmol})$, and allyltrichlorosilane (75 μ L, 0.47 mmol) at 0 °C. The reaction mixture was stirred at 20° C for 72 h. The reaction was quenched with saturated aqueous NaHCO₃ (1 mL), the layers were separated and dried over MgSO4. GC yield of 9 was 50%. Chiral GC (HP-Chiral β 30 m \times 0.25 mm, oven: 80 °C for 15 min, then $1 \degree$ C/min to $150 \degree$ C, 5 min at that temperature) showed 20% ee (t_R = 57.55 min, t_S = 56.03 min).

4.3. Enantioselective alkylation of benzaldehyde with diethylzinc to $(R)-(+)$ -1-phenyl-1-propanol (11)

To a solution of $(R)-(+)$ -6 (7 mg, 0.02 mmol) in toluene (0.6 mL) 1 M solution of diethyl zinc hexane (0.68 mL, 0.68 mmol) was added at 0° C and the reaction mixture was stirred for 20 min. Then benzaldehyde $(33 \mu L, 0.33 \text{ mmol})$, was added and the reaction mixture was stirred for 72 h at 20 °C. The reaction was quenched with 10% H₂SO₄ (0.5 mL), the organic layer was separated and the aqueous phase was extracted with diethylether, the combined organic layers were dried over $MgSO₄$. GC yield of the 1-phenyl-1-propanol was 62%. Chiral GC (HP-Chiral β 30 m \times 0.25 mm, oven: 80 °C for 15 min, then 1 \degree C/min to 150 \degree C, 5 min at that temperature) showed 17% ee (t_R =49.78 min, t_S =51.09 min).

4.4. Crystallography

Crystal data for: 3g. $C_{23}H_{23}NO$, $M=329.42$, monoclinic, $P2_1/n$, $a=13.1910(3)$ Å, $b=8.6090(2)$ Å, $c=16.4300(4)$ Å, $\beta=107.1860(11)^\circ$, $V=1782.50(7)$ \AA^3 , $Z=4$, $D_x=$ 1.228 Mg m⁻³. A colorless prism dimensions $0.5 \times 0.25 \times$ 0.2 mm was mounted on glass capillary with epoxy glue and measured at Nonius KappaCCD diffractometer by monochromatized Mo K α radiation (λ =0.71073 Å) at 150(2) K. An absorption was neglected $(\mu = 0.074 \text{ mm}^{-1})$; a total of 26146 measured reflections in the range $h=-17$ to 17, $k=-11$ to 11, $l=-21$ to 21 ($\theta_{\text{max}}=27.5^{\circ}$), from which 4076 were unique (R_{int} =0.015), 3294 observed according to the $I>2\sigma(I)$ criterion. The structure was solved by direct methods $(SIR92)^{32}$ $(SIR92)^{32}$ $(SIR92)^{32}$ and refined by full-matrix least squares based on F^2 (SHELXL97).^{[33](#page-138-0)} The hydrogen atoms

were recalculated into idealised positions (riding model) and assigned displacement parameter either $H_{\text{iso}}(H)$ = 1.2 U_{eq}(pivot atom) or $H_{iso}(H) = 1.5$ U_{eq}(pivot atom) for methyl moiety. The refinement converged $(\Delta/\sigma_{\text{max}}=0.001)$ to $R=0.0447$ for observed reflections and $wR=0.124$, S= 1.038 for 229 parameters and all 4076 reflections. The final difference map displayed no peaks of chemical significance $(\Delta \rho_{\text{max}} = 0.252, \ \Delta \rho_{\text{min}}^{\text{1}} - 0.246 \ \text{eA}^{-3}).$

Crystal data for $(R)-(+)$ -6 \cdot (S)-(-)-7: C₂₃H₂₃NO₂ \cdot C₂₀- $H_{14}O_2$, $M=631.74$, monoclinic, P_{21} , $a=11.5900(3)$ Å, $b=9.0280(2)$ Å, $c=16.0570(4)$ Å, $\beta=104.4160(13)^\circ$, $V=$ 1627.22(7) \mathring{A}^3 , $Z=2$, $D_x=1.289$ Mg m⁻³. A colorless plate of dimensions $0.37 \times 0.2 \times 0.15$ mm was mounted on glass capillary with epoxy glue and measured at Nonius KappaCCD diffractometer by monochromatized Mo K α radiation ($\lambda = 0.71073$ Å) at 150(2) K. An absorption was neglected $(\mu = 0.082 \text{ mm}^{-1})$; a total of 19962 measured reflections in the range $h=-15$ to 15, $k=-11$ to 11, $l=-20$ to 20 ($\theta_{\text{max}}=27.5^{\circ}$), from which 7408 were unique (R_{int} =0.028), 6678 observed according to the $I>2\sigma(\hat{I})$ criterion. The structure was solved by direct methods $(SIR92)^{32}$ $(SIR92)^{32}$ $(SIR92)^{32}$ and refined by full-matrix least squares based on F^2 (SHELXL97).^{[33](#page-138-0)} The absolute configuration of the crystal has been assigned by reference of known configuration of (S) - $(-)$ -binol 7. The hydrogen atoms were found on difference Fourier map, those on carbon atoms were recalculated into idealised positions (riding model) and assigned displacement parameter either $H_{iso}(H) = 1.2$ U_{eq}(pivot atom) or $H_{\text{iso}}(H) = 1.5 \text{ U}_{\text{eq}}(\text{pivot atom})$ for methyl moiety. The hydrogen of the hydroxyl moiety was refined isotropically. The refinement converged ($\Delta/\sigma_{\text{max}}$ =0.000) to R= 0.0378 for observed reflections and $wR = 0.0956$, S 1.001 for 443 parameters and all 7408 reflections. The final difference map displayed no peaks of chemical significance $(\Delta \rho_{\text{max}} = 0.259, \ \Delta \rho_{\text{min}} - 0.181 \text{ eA}^{-3}).$

Crystallographic data for structural analysis have been deposited with the Cambridge Crystallographic Data Centre, CCDC no. 283868 and 283869 for 3g and $(R)-(+)$ -6 $(·)$. $(-)$ -7, respectively. Copies of this information may be obtained free of charge from The Director, CCDC, 12 Union Road, Cambridge CB2 1EY, UK (fax: $+44$ 1223 336033; e-mail: deposit@ccdc.cam.ac.uk or [www: http://www.ccdc.](http://www:http://www.ccdc.cam.ac.uk) [cam.ac.uk](http://www:http://www.ccdc.cam.ac.uk)).

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.tet.2005.10.](http://dx.doi.org/doi:10.1016/j.tet.2005.10.034) [034.](http://dx.doi.org/doi:10.1016/j.tet.2005.10.034) Preparation and spectral characteristics of the starting 1-aryl-1,7-diynes 2 and the pyridine 4.

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Synthesis and characterization of chiral, bridged resorcinarenes as templates for asymmetric catalysis

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Abstract—A full study of the synthesis of chiral, bridged resorcinarenes (3a–3l, 13a, 13b) is presented using Mannich condensation of C_{2v} tetraprotected resorcinarenes with chiral $1, n$ -diamines bearing homochiral α -methylbenzyl auxiliaries at each terminal nitrogen. The study has revealed the methodology to be applicable to preparing a broad range of bridged structures with varying lengths of bridge, different functionality in the bridge and various protecting groups on the upper rim. Reproducible and satisfactory yields in the reaction were only obtained with the pendant R group as methyl. The bridged adducts have been fully characterized by a range of spectroscopic techniques, and NMR has revealed varying trends in the way the various bridges protrude into the cavity. Low temperature NMR as well as X-ray structures of tetramesylate 15 and tetratoluate 3g has revealed hydrogen bonding to the amine nitrogens in the bridge to be an important control element for positioning the bridge relative to the cavity of the bowl. The derivatives provide chiral templates for asymmetric catalysis studies using cooperative effects in the bowl.

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1. Introduction

It has long been a desire for synthetic chemists to achieve in the laboratory what is seemingly so effortlessly achieved in biological systems. While it is well known that enzymes can efficiently regio- and stereoselectively catalyse a vast array of organic transformations under mild conditions, a synthetic example has yet to be produced that can match an enzyme in rate acceleration, turnover and specificity.^{[1](#page-152-0)} Recently, attention has been directed at utilizing calixarenes and resorcinarenes as platforms for asymmetric processes with many examples now reported on the use of these frameworks for chiral recognition and discrimination studies.^{[2](#page-152-0)} Regarding catalysis, while the literature contains examples of calixarenes and resorcinarenes that are used as catalysts,^{[3](#page-152-0)} few of them describe asymmetric catalysis. In this regard, Matt has shown that a lower-rim, inherently chiral calixarene scaffold can be used in allylic alkylation (palladium) and hydrogenation (rhodium), although low ees were obtained.^{[4](#page-152-0)} Others have shown that cooperative effects may potentially be provided by supramolecular interactions involving the concave bowl.^{[5](#page-152-0)} However, a comprehensive picture has yet to emerge regarding application of directing effects in the bowl towards designing superior asymmetric catalysts for carrying out reactions rather than purely for

recognition phenomenon. We have recently reported the first example of an asymmetric reaction taking place in the bowl,^{[6](#page-152-0)} which we have termed intracavity asymmetric catalysis, using chiral, distally-bridged resorcinarenes of the type shown in Figure 1. In this paper we discuss the scope of the methodology used for the synthesis of chiral bridged resorcinarenes, along with new examples for use in the study of asymmetric reactions occurring in the bowl.

Figure 1. Example of a chiral, distally-bridged resorcinarene.

2. Strategy

Our strategy ([Scheme 1\)](#page-140-0) involved using the Mannich reaction to prepare chiral, bridged resorcinarenes^{[6](#page-152-0)} as C_{2v} symmetric templates for catalysis, in which a functional group in the middle of the bridge would offer opportunities for intracavity catalysis. Our choice of Mannich

Keywords: Resorcinarene; Catalytic template; Mannich reaction.

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Scheme 1. Retrosynthetic strategy for formation of bridged resorcinarenes.

methodology for bridge introduction was inspired by a precedent set by Böhmer and Shivanyuk,^{[7,8](#page-152-0)} who demonstrated that primary diamines undergo Mannich reactions with C_{2v} –tetraprotected resorcinarenes and formaldehyde to afford bridged bis-benzoxazines. We sought to adapt this precedent to using secondary 1,n-diamines bearing a chiral a-methylbenzyl unit at each nitrogen as a chiral auxiliary for generation of enantiomerically pure C_{2v} -symmetric bridged resorcinarenes as catalyst templates. This methodology would provide the opportunity to incorporate desirable functionality in the bridge by using functionalized diamine lines of type 1 (varying n and X). In addition, other structural parameters for developing a structure–activity profile of catalysis included the choice of upper-rim protecting groups Y in the C_{2v} -array of resorcinarene 2, as well as the pendant benzylic R groups.

Our previous work^{[6](#page-152-0)} described the synthesis of bridged resorcinarenes with Y = tosylate, $n=1$ and varying length of benzylic R with or without ketal X groups in the bridge as described in Table 1. In the comprehensive study described in this paper, emphasis was focused on two parameters, namely the length and nature of the bridge $(n \text{ and } X)$ and the size and nature of the upper-rim protecting groups (Y).

Table 1. Previously synthesized chiral bridged resorcinarenes

		R	n	X, X
3a	Tos	CH ₃		Н
3 _b	Tos	C_5H_{11}		Н
3c	Tos	$C_{11}H_{23}$		Н
3d	Tos	CH ₃		OMe
3e	Tos	CH ₃		$OCH2CCH3)2CH2O$

In order to install different upper-rim protecting groups on the bridged resorcinarenes, it was necessary to synthesize a series of appropriately C_{2v} -protected resorcinarenes, which could be used in the Mannich reaction with suitable diamines (cf. Scheme 1). This could be achieved by using and expounding methodology developed by Shivanyuk and Böhmer in 1[9](#page-152-0)98,⁹ involving either direct functionalisation with 4 equiv of electrophile or through formation of the tetra-Cbz derivative, which may then be used to form other C_{2v} -protected resorcinarenes via a protection, deprotection (CBz) protocol. The latter methodology is powerful in view of its

practical simplicity, since the desired $C_{2\nu}$ -tetra-Cbz derivative intermediate precipitates out of the reaction medium as a complex with 2 equiv triethylammonium hydrochloride, thus ensuring high purity and easy access to the desired product. After functionalising the remaining four hydroxyl groups with a suitable electrophile, the Cbz groups can be removed by facile hydrogenolysis furnishing new C_{2v} -tetraprotected resorcinarenes.

There were many possible protecting-group targets for our study, but essentially the aim was to evaluate a selection of both large and small ones that might affect the steric environment in the bowl of the final catalyst. One restriction in setting about these syntheses was that the methodology was only really practical on the tetramethylresorcinarene (2) ; $R=CH_3$, $Y=H$), since resorcinarenes with longer pendant chains generally gave little or no product in the acylation reactions.^{[9](#page-152-0)} Further studies in this area, though, may result in C_{2v} -protected resorcinarenes with longer pendant chains such as the tetraundecylresorcinarene tetratosylate (2c; $R = C_{11}H_{23}$, Y = Tos) used to synthesize 3c.^{[10](#page-153-0)}

The large number of possibilities for protecting groups resulted in the decision to evaluate other sulfonates of varying sizes, in view of the success of the tetratosylates. A methanesulfonyl group (Mes) was identified as a good example of a smaller protecting group than tosylate, with 2,4,6-triisopropylbenzenesulfonate (Trips) as one with greater steric bulk. It was also thought desirable to evaluate groups other than the sulfonate functionality in order to probe electronic effects. Thus the bridged resorcinarene tetrabenzyloxycarbonate (Cbz) was identified, also in view of its ready availability^{[9](#page-152-0)} as already mentioned, as well as the resorcinarene tetra $(p$ -tolyl) (Tol) ester since it was also available by direct acylation of the parent resorcinarene with *p*-toluoyl chloride.^{[9](#page-152-0)} It was thought that evaluating this bridged analogue would provide an interesting comparison to the resorcinarene tetratosylate 3a in view of similarity in steric size but difference in donor character around the respective oxygen atoms. Likewise, the resorcinarene tetraacetate (Ac) was also considered as one for making an informative comparison with the resorcinarene tetramesylate on steric grounds while at the same time complementing the ester series. A tetramethylated resorcinarene was also targeted for this study.

Scheme 2. Direct synthesis of C_{2v} -protected resorcinarenes. Reagents and conditions: (i) YCl (4 equiv), Et₃N (4 equiv), CH₃CN, rt.

3. Results and discussion

3.1. Synthesis of alternative C_{2v} -protected resorcinarenes

Formation of the sterically demanding C_{2v} -symmetrical resorcinarene tetratripsylate 2f (2,4,6-triisopropyl-benzenesulfonate) was gratifyingly achieved in 34% yield[†] using the methodology developed by Shivanyuk for tetratosylation (Scheme 2).¹¹ Similarly, both the C_{2v} -resorcinarene tetratoluate 2g as well as the all-important C_{2v} -resorcinarene tetrabenzyloxycarbonate (CBz) 2h could be readily obtained via the said literature procedure in 21 and 41% yields, respectively.^{[9](#page-152-0)}

The resorcinarene tetraacetate 2k could be synthesized by the method reported by Shivanyuk, involving acetylation of the tetracarbonate 2h to give 2i in excellent yield followed by quantitative cleavage of the protecting groups.^{[9](#page-152-0)} For this reaction, the authors used catalytic hydrogenolysis over palladium supported on carbon in dioxane as a solvent. This was probably due to the compound's poor solubility in ethanol. In the present work it was found that performing the reaction in a solvent mixture of ethanol and THF (1:1 v:v) at room temperature gave a better reaction profile than that from using dioxane. Formation of the resorcinarene tetramesylate 2l was also achieved in the same way as the resorcinarene tetraacetate 2k [\(Scheme 3](#page-142-0)). To this end, mesyl chloride was reacted with resorcinarene tetracarbonate 2h in the presence of Hünig's base to give the required tetramesylate derivative 2j in a good yield (75% after recrystallization of the crude reaction product). The mesylation was rapid being finished by tlc within 30 min (from 0° C to rt). Removal of the carbonate groups could likewise be achieved as before using catalytic hydrogenolysis. Once again, a 1:1 mixture of ethanol and THF was the preferred solvent, cleanly affording the desired tetramesylate 2l after 6 h at room temperature.

It was also thought desirable to tetramethylate the resorcinarene. Though resorcinarene tetratosylate 2a could be tetramethylated using dimethyl sulphate, the tosyl groups failed to be cleaved under basic conditions, a fact already noted by Shivanyuk.^{[9](#page-152-0)} Tetramethylation of the resorcinarene tetracarbonate 2h or tetratoluate 2g unfortunately failed to give the desired C_{2v} -symmetrical product. Many conditions were tried, using both dimethyl sulfate or methyl iodide as methylating agent, and varying the type of base used (sodium hydride, triethylamine, Hünig's Base). In one case the reaction was attempted in neat methyl iodide, but in all attempts the reactions gave mixtures of 3–5 unidentified products, which were inseparable by chromatography. It became clear, after stirring the tetracarbonate with sodium hydride that the failure of these reactions was due to the starting material undergoing some manner of transformation. Tentatively it could be suggested that this was due to the carbonate groups migrating around the resorcinarene ring. Unfortunately, the phenolic hydroxyl oxygens are not nucleophilic enough for direct alkylation and thus require the use of a base to form the phenoxide, which resulted in starting-material transformation. Direct methylation of tetramethylresorcinarene (2; $R=CH_3$ and Y=H) did not give the desired product; one isolated product revealed tetramethylation by ¹H NMR but did not show $C_{2\nu}$ symmetry.

3.2. Synthesis of bridged resorcinarenes

The synthesis of bridged resorcinarenes containing different protecting groups, $n=1$ and $X=H$, was achieved using the methodology already described for synthesizing the bridged resorcinarene tetratosylates.^{[10](#page-153-0)} Thus each C_{2v} -symmetrical tetraprotected resorcinarene 2a–2l was heated in acetonitrile with a slight excess of the diamine and paraformaldehyde in large excess to yield distally-bridged products 3a–3l ([Scheme 4](#page-142-0)). Such conditions were found to be superior to reaction with aqueous formaldehyde in ethanol, which resulted in resorcinarene ethoxymethylation. Yields were generally good except for the resorcinarene tetraacetate, which was not optimized ([Table 2\)](#page-142-0).

[†] Yields for these types of reactions are typically less than 50% even dropping to less than 10% owing to the formation of isomeric products.

Scheme 3. Reagents on conditions: (i) XCl (5 equiv), i -Pr₂NEt (6 equiv), DCM, rt; (ii) H₂, Pd/C (10%), TH–FEtOH (1:1), rt.

Scheme 4. Reagents and conditions: (i) diamine 1 (1.2 equiv), $(CH_2O)_n$ (10 equiv), 90 °C.

Table 2. Results for bridging reactions

	Y	Yield $(\%)$
3a	Tos	60
3f	Trips	78
$\frac{3g}{3h}$	Tol	55
	Cbz	70
3k	Ac Mes	20
3 _l		51

3.3. Synthesis of longer diamines

The second parameter evaluated was the synthesis of bridged resorcinarenes with longer bridges. Therefore it became necessary to synthesize a longer diamine than type 1. This could be achieved using pimelic acid (heptanedioic acid) 4 as starting material as shown in Scheme 5. Thus pimelic acid was readily coupled to α -methylbenzylamine under DCC conditions yielding the desired diamide 5 in excellent yield. In this reaction the addition of hydroxybenzotriazole (HOBt) was found to be important for preventing the formation of the N-acylated urea sideproduct. Reduction of the diamide was achieved using lithium aluminium hydride in refluxing THF to give diamine 6 in 95% yield for each step.

In a similar fashion, synthesis of the dimethoxy ketalfunctionalized diamine required 4-oxopimelic acid, which could be derived from furfural in three steps. The first of the steps converted furfural 7 into 3-(2-furyl)acrylic acid 8 via a Knoevenagel reaction, 12 by heating furfural with malonic acid in pyridine at 100 °C for 2 1/2 h to give 8 in a good overall yield (69% after recrystallization). Conversion of acid 8 into the desired oxopimelic acid 9 was efficiently carried out in two further steps by first refluxing the acid 8 in ethanol using a continuous passage of HCl gas. 13 13 13 Distillation of the crude material obtained gave diethyl pimelate as a clear oil which was readily saponified with aqueous potassium hydroxide (3 M) at room temperature in under 30 min, resulting in 4-oxopimelic acid 9 in 72% overall yield. The coupling of α -methylbenzylamine was again accomplished via a DCC coupling with the diacid to give diamide 10 in good yield (83%). The penultimate step required protection of the ketone carbonyl group as its dimethoxy ketal, prior to reduction of the diamide. This was

Scheme 5. Reagents and conditions: (i) α -methylbenzylamine, DCC, HOBt, CH₂Cl₂; (ii) LiAlH₄, THF, reflux.

Scheme 6. Reagents and conditions: (i) malonic acid, pyridine, 100 °C; (ii) EtOH, HCl, reflux; (iii) KOH, H₂O, rt; (iv) α -methylbenzylamine, DCC, HOBt, CH_2Cl_2 , rt; (v) $CH(OCH_3)_3$, MeOH, p-TsOH (cat), reflux; (vi) LiAlH₄, THF, reflux.

achieved by refluxing 10 in excess trimethyl orthoformate and catalytic p-toluenesulfonic acid overnight. This cleanly afforded the desired ketal 11 in yields of up to 92%. Finally, reduction of diamide 11 to diamine 12 was found to be possible using lithium aluminium hydride in refluxing diethyl ether, with the addition of triethylamine to attenuate the slight Lewis acidity of the metal salts, Scheme 6.

3.4. Synthesis of longer bridged resorcinarenes

The newly prepared longer chiral diamine line 6 was used to bridge resorcinarene tetratosylate 2 (Y=Tos, R=CH₃). Using the standard reaction conditions, a maximum yield of 36% of the bridged product 13a was obtained (Scheme 7). The low yield of bridged resorcinarene 13a was in contrast to the better yields that were obtained for the bridged resorcinarene using the shorter diamine line 3a.

This lower yield deserves mechanistic comment. It is reasonable to assume that the bridging process is achieved via two discrete steps, the first being the initial aminomethylation, which is followed by a second, rate-determining aminomethylation step to close the bridge. The lower yield for 13a from the longer line 6 may be accounted for by assuming closure to be slower than for 3a on entropic grounds (larger $\Delta S^{\#}$ -ve). The longer lifetime for the mono-adduct from 6, would then increase the chances of an intermolecular reaction. Careful scrutiny of the reaction tlc revealed another product, which was slightly more polar than the bridged product 13a. It had virtually identical

¹H and ¹³C NMR spectra to the bridged product $13a$, the only difference being that the signals in the 1 H NMR were a little less resolved. Its IR spectrum was no different to 13a, but its optical rotation was opposite in sign and different in magnitude $(-7.0^{\circ}$ vs $+13.2^{\circ}$ for 13a). MALDI-TOF mass spectrometry revealed that this compound had a molecular mass of 3047 amu corresponding to exactly twice the molecular mass of bridged resorcinarene 13a. This prompted us to conclude that the side-product isolated in 5% yield was dimer 14 [\(Fig. 2\)](#page-144-0).

This structure seems plausible as similar examples exist in the literature, which have been well-documented and reviewed.^{[14](#page-153-0)} In 1998, Böhmer demonstrated this type of dimerisation with tetrapentylresorcinarene 2 ($Y = H$, R= C_5H_{11}) and ethylenediamine under Mannich conditions, giving an octabenzoxazine dimer linked by four bridges.^{[15](#page-153-0)} Böhmer reported that only ethylenediamine worked, whereas 1,3-diaminopropane, and ethylenediamine derivatives such as $1,2$ -diaminocyclohexane or N, N' -dimethylethylenediamine failed to give any defined reaction product.[15](#page-153-0) This chiral resorcinarene dimer linked by two bridges represents the first example of a dimer of this type.

In a similar fashion, the functionalized, longer ketal-diamine 12 could be used to bridge resorcinarene 2 ($Y = T\text{cos}$, $R = CH₃$). This was performed in the usual manner at 90 °C and gave 10% of the desired product 13b after one hour. Increasing the number of equivalents of diamine from one to three gave a vastly improved yield of 48%. With a slight

Figure 2. Proposed structure of dimer 14.

decrease in reaction temperature, the yield could be further optimized to 57%. Interestingly, in this case no dimer like 14 was observed, presumably because of the steric presence of the ketal in the middle of the line.

4. Spectroscopic and conformational aspects

The NMR spectra of the bridged resorcinarenes were remarkably straightforward to assign, owing to the high degree of symmetry; a typical example of a ${}^{1}H$ NMR spectrum is shown in Figure 3 for resorcinarene 3h $(Y = CBz)$. All of the NMR signals could be unambiguously assigned using 2D-NMR techniques (HSQC and HMBC). A few significant features of the ¹H NMR spectra deserve comment. Firstly, the three singlets (marked with *) for the resorcinarene aromatic methine protons of the bridged products were used as markers to indicate that reaction had successfully taken place at the positions *ortho* to both hydroxyl groups of the tetraprotected resorcinarene.

Secondly, a new signal between 3.71 and 3.78 ppm (singlet) in Figure 3, although in some cases appearing as a pair of doublets owing to diastereotopicity; marked with \downarrow) was always observed for these structures corresponding to the methylene protons on the carbon attaching the bridge to the

resorcinarene. Thirdly, no hydroxyl signals were observed suggesting a fast interconversion between hydrogen-bonded modes to be occurring that creates very broad peaks on the timescale of the NMR experiment. Fourthly, there was an upfield shift of all the methylene protons on the bridge relative to the starting diamine, with the central methylenes experiencing the greatest shift; this feature is an important indication that the bridge resides over the bowl, resulting in anisotropic shielding by the aromatic rings comprising the bowl.[16](#page-153-0) An increase in upfield shift from one resorcinarene to another was interpreted as indication of the bridge spending more time deeper within the cavity of the bowl. Table 3 shows the variation in the chemical shift of the central methylene group for the bridged resorcinarenes synthesized. Going from 3a to 3b and then to 3c involving lengthening of the pendant alkyl chain creates a conformational change that encourages the bridge to spend more time lower within the cavity. This may be due to increased repulsion between the pendant R groups, resulting in the upper-rim aromatic rings to fold inwards slightly forcing the bridge to bury itself more into the cavity. In the case of 3a versus 3h it would appear that the CBz protecting groups have a similar effect, also resulting in the bridge to lie lower in the cavity. Conversely, the sterically undemanding mesyl groups with CH_3 as the pendant R group (3l) releases the line out of the cavity as indicated by a weaker shielding effect compared to 3a. In the case of the longer bridge 13a, the central methylene protons do not exhibit as great an upfield chemical shift as 3a, indicating that the longer bridge prefers to sit higher above the cavity than deeper in it, presumably to maximize entropy.

Table 3. Comparison of chemical shifts for central methylene protons

	n	Y	R	Central methylene (ppm)	Central methylene Δ ppm
1				1.27	
3a	1	Tos	CH ₃	0.90	-0.37
3 _b		Tos	C_5H_{11}	0.72	-0.55
3c		Tos	$C_{11}H_{23}$	0.70	-0.57
3f		Trips	CH ₃	0.99	-0.28
3g		Tol	CH ₃	0.93	-0.34
3h		Cbz	CH ₃	0.77	-0.50
3k		Ac	CH ₃	0.84	-0.43
31		Ms	CH ₃	1.05	-0.22
6	2			1.32	
13a	\overline{c}	Tos	CH3	1.10	-0.17

Figure 3. 1 H NMR spectrum of resorcinarene 3h.

Figure 5. High and low temperature ${}^{1}H$ NMR spectra of 3a.

Finally, another important observation was the downfield shift (> 0.5 ppm) of the protons on the lower rim of the protected resorcinol rings. Prior to bridging they were found around 6.25 ppm, significantly upfield to what would be expected. The reason for this has been attributed to the resorcinarene existing in a boat conformation, 17 since the aromatic groups that are most horizontal have their protons on the lower rim in the shielding zone of the vertical aromatic rings. In the bridged compounds, the protons appeared downfield at around 6.85 ppm $(+/-0.20$ ppm), suggesting a change in shape towards the crown conformation, though to what extent this change had occurred was not pursued any further by NMR studies.

The possibility of hydrogen bonding between the tertiary amine nitrogens on the bridge and the phenolic hydroxyl groups on the resorcinarene was also considered as an important structural feature. Such bonding would generate three diastereomeric structures as shown in Figure 4; two with C_2 symmetry and one with C_1 symmetry. The C_1 symmetric structure would be expected to show 5 singlets for its aromatic methine protons in its ${}^{1}H$ NMR spectrum, while the C_2 -symmetric isomers would show three singlets each. This distribution as expected was not seen at temperatures above room temperature, owing to conformational exchange. However, on cooling to $0^{\circ}C$ the resorcinarene aromatic signals became very broad and indistinct. Further cooling to -60° C revealed a far more complex picture, in which the rate of 'flipping' between the

different hydrogen bonded modes slowed enough for the individual diastereomers in Figure 4 to be observed. Figure 5 shows the aromatic region of the ${}^{1}H$ NMR spectrum of 3a at 50 and -60 °C. At 50 °C there are 3 signals (marked with \downarrow) at 6.79, 6.83 and 7.11 ppm for the average of the conformers. On cooling, the signals become increasingly broadened and then resolved below -40 °C into separate signals, becoming clear at -60° C. By careful inspection and integration it was possible to assign the signals for the C₁ conformer (5 signals marked with \bullet), while 2 sets of 3 signals each could be seen for the C_2 conformers (marked with) One extra signal could be attributed to a hydroxyl proton.‡ Integration revealed that the different conformers existed in an approximate ratio of 1:1:1 $(C_2:C_2:C_1)$. The signal at δ = 6.92 ppm integrates for 2H.

It is possible that H-bonding could affect the conformation of the bridged resorcinarene, and therefore the hydroxyl substituents were suitably protected. The latter are however not easy to functionalize, 18 and it was thus decided to use mesyl chloride as a small reactive electrophile. Mesylation using excess mesyl chloride and Hünig's base in dichloromethane was successful in furnishing the desired tetramesylated product 15 in a favourable 87% yield [\(Scheme 8\)](#page-146-0).

[‡] As mentioned, the hydroxyl protons are invisible at normal NMR temperatures owing to the rapid hydrogen bonding exchange. On cooling, the signals at 6.24 ppm and at 13.10 ppm (not shown) become visible.

Scheme 8. Reagents and conditions: (i) mesyl chloride (20 equiv), i-Pr₂EtN (20 equiv), DCM, rt.

Figure 6. 1 H NMR spectrum of resorcinarene 15.

The ${}^{1}H$ NMR spectrum (Fig. 6) of 15 provided some surprises, notably that the methylene protons in the centre of the bridge were now found at -0.95 ppm, representing a massive 2.22 ppm upfield shift from the diamine 1 (prior to bridging). This could only be attributed to the bridge spending a much higher percentage of its time deep within the resorcinarene cavity. This might arise from the disruption of the hydrogen bonding with the amines, allowing the bridge to 'collapse' into the cavity, analogous to what happens if the supports of a suspension bridge are removed.

Furthermore, another intriguing spectroscopic observation was a 0.45 ppm upfield shift of the signals for the pendant methyl groups on the resorcinarene (from 1.37 to 0.89 and 0.93 ppm). The reason for this was not made clear until a single crystal X-ray structure determination was realised. Great difficulty was experienced in trying to grow suitable crystals for X-ray diffraction. Eventually a suitable crystal was found in a solution of toluene, acetone and dichloromethane. 15 crystallized in the space group $P2_12_12$ with $Z=4$, with two dichloromethane molecules of solvation which were located in five different positions in the unit cell with partial occupancies. As a result, the structure could not be refined satisfactorily because the dichloromethane molecules were highly disordered with large temperature factors. Therefore, the fractional atomic coordinates of the structure are not reported. Only the unit cell parameters are listed in the supplementary data. The positions of the resorcinarene atoms alone were located unambiguously and its molecular conformation can thus be reported. This is given by the torsion angles listed in the supplementary data.

The X-ray picture of 15 as an ORTEP diagram (ellipsoidal model at 30% probability level) revealed the bridge to be positioned inside the cavity in accordance with predictions

from the 1 H NMR, and also provided the answer to the upfield shift of the pendant methyl groups. As can be seen in Figure 7, the aromatic groups of the tosylate groups are pointing downwards towards the lower rim in order to

Figure 7. X-ray structure of 15.

Figure 8. X-ray structure of 3g.

minimize repulsion with the mesylate groups. As a result, they are in a position to anisotropically shield the pendant methyl groups.

It is clear from this X-ray structure that the hydrogen bonding in the bridged resorcinarene is important in maintaining a preferred shape by: (a) keeping the bridge above the cavity, potentially allowing reactions to be accommodated there; and (b) holding the protecting groups up instead of down allowing for potential cooperative effects in the bowl.

It was thus hoped that an X-ray structure of a hydrogenbonded bridged resorcinarene could be obtained. Suitable crystals were not readily obtainable, but eventually slow evaporation of a solution of tetratoluate 3g in diisopropyl ether, pentane and dichloromethane resulted in the formation of diffraction-quality crystals, although they were very unstable outside their mother liquor. Bridged resorcinarene 3g crystallized in the space group P1 with $Z=2$ as a dipentane solvate. There are thus two independent resorcinarene host molecules and four pentane guest molecules in the unit cell, all located at general positions. We note that the comformation of the two independent resorcinarene molecules are very similar, and Figure 8 shows one of these structures. The unit cell parameters and other crystal data and refinement data are given in the supplementary data.

The X-ray structure of 3g reveals a boat-like conformation and demonstrates the effect that hydrogen bonding has on the positioning of the bridge, sustaining it over the cavity and with the protecting toluate ester groups held up towards the cavity (as apposed to hanging downwards as in 15).

5. Conclusions

This paper has extended the methodology developed previously, 6.9 to the synthesis of a range of new, chiral, bridged resorcinarenes, which have been fully characterized by NMR and X-ray techniques resulting in the observation of some interesting aspects of bridge-positioning relative to the cavity of the bowl. In the following paper, a comprehensive study is presented on the use of these derivatives as asymmetric templates for the catalyzed enantioselective addition reaction of diethylzinc to benzaldehyde, which is considered to be one the first reported examples of using the bowl of an asymmetrically functionalized resorcinarene for promoting asymmetric catalysis,^{[6](#page-152-0)} as suggested recently by Iwanek.^{[19](#page-153-0)} The results are rationalized using a stereoselectivity model.

6. Experimental

6.1. General remarks

All reactions were carried out under nitrogen using dry solvents. Nuclear Magnetic Resonance spectra were recorded on a Varian Unity 400 (100 MHz for 13 C) or Varian Mercury 300 MHz (75 MHz for ¹³C) and were carried out in chloroform-d. Optical rotations were obtained using a Perkin Elmer 141 polarimeter at 20 °C. Melting points were obtained using a Reichert Jung Thermovar hotstage microscope and are uncorrected. Elemental analyses were performed using a Fisons EA 1108 CHN elemental analyser. Infrared spectra were recorded on a Perkin–Elmer Paragon 1000 FT-IR spectrometer in either dichloromethane or chloroform. Resorcinarenes 2a,g–i were synthesized according to the literature procedures.^{[9](#page-152-0)} Bridged resorcinarenes $3a-e$ have been reported previously.^{[6,10](#page-152-0)} Crystallographic data for the structures in this paper have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication numbers CCDC 273318 and 273319. Copies of the data can be obtained free of charge from CCDC via [www.ccdc.cam.ac.uk/](http://www.ccdc.cam.ac.uk/data_request/cif) [data_request/cif.](http://www.ccdc.cam.ac.uk/data_request/cif) Names and numbering of compounds follows IUPAC 'phane' nomenclature rules;[§] [Figure 9](#page-148-0) shows two examples of this.

6.1.1. $3^4, 3^6, 7^4, 7^6$ -Tetrahydroxy-2,4,6,8-tetramethyl-1⁴,1⁶,5⁴,5⁶-tetra(2',4',6'-triisopropylbenzene-sulfonyloxy)-1,3,5,7(1,3)tetrabenzenacyclooctaphane (2f). Tetramethyl-resorcinarene 2 ($R=CH_3$, $Y=H$) (3.27 g, 6 mmol) was dissolved in acetonitrile (60 mL) and triethylamine (3.35 mL, 24 mmol) added. The pink, heterogeneous solution was stirred at room temperature for 15 min and then 2,4,6-triisopropylbenzenesulfonyl chloride (5.16 g, 6 mmol) was added in one portion. The reaction was allowed to stir at room temperature overnight, followed by filtration of the solid precipitate formed. The precipitate was washed with cold acetonitrile $(2 \times 20 \text{ mL})$ and then dissolved in dichloromethane (75 mL). 1 M HCl (100 mL) was added and the product extracted into dichloromethane $(3 \times 50 \text{ mL})$. After drying (MgSO₄) and evaporation, the crude product was further purified by column

[§] For a full description see <http://www.chem.qmul.ac.uk/iupac/>

Figure 9. Example of numbering format for resorcinarenes.

chromatography (60 g silica gel, eluting with ethyl acetate/ petroleum ether 1:4). In this way, tetratripsylresorcinarene $2f(3.3 g, 34\%)$ could be prepared as a white solid. Mp 190– 192 °C (from DMF/ethyl acetate/petroleum ether); $v_{\text{max}}/$ cm^{-1} (CHCl₃) 3476s + 3387s (O–H, H-bonded), 3015s (C–H, aromatic), $2964s + 2931$ (CH₃), $1599m + 1514s$ (aryl stretch), $1346s + 1180s$ (–SO₂–); δ_H (300 MHz, CDCl₃) $1.11 + 1.13$ (48H, 2 \times d, $J = 6.8$ Hz, $7^{7}/9^{7} - i$ -Pr), 1.21 (24H, d, $J=6.8$ Hz, $8'-i$ -Pr), 1.38 (12H, d, $J=6.8$ Hz, $-CH_3$), 2.88 $(4H, spt, J=6.8 Hz, H-8', 3.93 (8H, spt, J=6.8 Hz, H-7'/9'),$ 4.39 (4H, q, J=6.8 Hz, H-2,4,6,8), 6.02 (2H, s, H-1⁵,5⁵), 6.31 (2H, s, H-3²,7²), 6.57 (2H, s, H-3⁵,7⁵), 7.03 (2H, s, H-1²,5²), 7.16 (4H, br, -OH), 7.18 (8H, s, H-3',5'); δ_C (75 MHz, CDCl₃) 20.0 (CH₃), 23.4 (8'*i*-Pr), 24.4 (7'/9'*i*-Pr), 29.9 (C-7 $'$ /9'), 31.9 (C-2,4,6,8), 34.2 (C-8'), 102.0 (C-1⁵,5⁵), 113.5 $(C-3^5,7^5)$, 119.0 $(C-1^1,1^3,5^1,5^3)$, 124.1 $(C-3^1,5^1)$, 125.2 $(C-1^2,5^2)$, 127.0 $(C-3^2,7^2)$, 129.5 $(C-1^2)$, 139.1 $(C-3^1,3^3,7^1,7^3)$, 144.8 $(C-3^4,3^6,7^4,7^6)$, 151.2 $(C-2^1,6^1)$, 153.2 (C-1⁴,1⁶,5⁴,5⁶), 154.5 (C-4^{*'*}); MALDI TOF: m/z (rel. int.) 1609.4 [M⁺ - H] (81), 1342.91 [M⁺ - SO₂Ar] (69), 1076.09 $[M^+-SO_2Ar \times 2]$ (70), 809.27 $[M^+-SO_2Ar \times 3]$ (53), 542.06 $[M^+ - SO_2Ar \times 4]$ (45). Found: C, 68.58; H, 7.54; S, 7.71%; $C_{92}H_{120}O_{16}S_4$ requires C, 68.63; H, 7.51; S, 7.97%.

6.1.2. 3^4 , 3^6 , 7^4 , 7^6 -Tetraacetoxy- 1^4 , 1^6 , 5^4 , 5^6 -tetrahydroxy-2,4,6,8-tetramethyl-1,3,5,7(1,3)tetrabenzenacyclooctaphane $(2k)^9$. Resorcinarene 2i $(1 g, 0.8 mmol)$ was dissolved in a 1:1 solvent mixture of THF and ethanol (50 mL). 10% Palladium on carbon (100 mg, 0.1 mmol) was added and the solution stirred at room temperature under a positive pressure of hydrogen. After 4 h the reaction was checked for completion (tlc) and the catalyst filtered off through Celite^w, washing with dichloromethane (100 mL) . Evaporation of the solvent afforded tetraacetate 2k (570 mg, 100%) as a pale pink solid. The ¹H NMR spectrum corresponded to that published in the literature.^{[9](#page-152-0)} $\delta_{\rm H}$ $(300 \text{ MHz}, \text{ DMSO-}d_6)$ 1.39 (12H, d, J=7.1 Hz, CH₃), 2.11 (12H, s, OAc), 4.39 (4H, q, $J=7.1$ Hz), 6.30 (2H, s, Ar-H), 6.41 (2H, s, Ar-H), 6.58 (2H, s, Ar-H), 7.33 (2H, s, Ar-H), 8.82 (4H, br s, –OH).

6.1.3. 3^4 , 3^6 , 7^4 , 7^6 -Tetra(benzyloxycarbonyloxy)- 1^4 , 1^6 , 54 ,56 -tetra(methylsulfonyloxy)-2,4,6,8-tetramethyl-1,3, 5,7 (1,3)tetrabenzenacyclooctaphane (2j). Diisopropylamine (3.06 mL, 17.6 mmol) and resorcinarene tetra-Cbz 2h (3.46 g, 3.2 mmol) were added to dry dichloromethane (40 mL) and the solution cooled to 0° C. Mesyl chloride (1.24 mL, 16 mmol) was added slowly and the reaction allowed to warm to room temperature. After 30 min the reaction was checked for completion (tlc) and then poured into 1 M HCl (100 mL). The product was then extracted into dichloromethane $(3 \times 70 \text{ mL})$, dried with magnesium sulphate and the solvent evaporated in the usual way. Recrystallization of the crude product from DMF/ethyl acetate/petroleum ether gave the title compound 2j (3.34 g, 75%) as a white powder. Mp 246-248 °C (from DMF/ethyl acetate/petroleum ether); v_{max}/cm^{-1} (CHCl₃) 1763s (C=O, ester), $1375s + 1169s$ (–SO₂–); δ_H (300 MHz, CDCl₃) 1.50 (12H, d, J=7.1 Hz, –CH₃), 2.88 (12H, s, –SO₂CH₃), 4.65 $(4H, q, J=7.1 \text{ Hz}, H-2,4,6,8), 5.25+5.30 (8H, 2 \times d, J_{AB} =$ 12 Hz, H-2'), $6.19 + 7.26$ (4H, s, H-1²,5² and H-3²,7²), 7.21 + 7.23 (4H, s, H-1⁵,5⁵ and H-3⁵,7⁵), 7.34–7.47 (20H, m, Ph); δ_C (75 MHz, CDCl₃) 20.0 (-CH₃), 31.9 (C-2,4,6,8), 38.1 ($-SO_2CH3$), 70.7 (C-2[']), 116.1 + 116.4, 125.6, 126.6, 128.6, 128.7, 128.8 (Ph + C-1²,5² and C-3²,7² and 3⁵,7⁵ and C-1⁵,5⁵), 134.6 (Ph), 134.3 + 135.5 (C-1¹,1³,3¹,3³,5¹,5³, 7^1 , 7^3), $145.5 + 146.6$ (C- 3^4 , 3^6 , 7^4 , 7^6 and C- 1^4 , 1^6 , 5^4 , 5^6), 153.2 (C-1'); MALDI-TOF m/z (rel. int.) 1431.51 $[M+K]^+(28)$, 1415.20 $[M+Na]^+(100)$. Found: C, 58.81; H, 4.66; S, 9.05%; $C_{68}H_{64}O_{24}S_4$ requires C, 58.61; H, 4.63; S, 9.20%.

6.1.4. 3^4 , 3^6 , 7^4 , 7^6 -Tetrahydroxy- 1^4 , 1^6 , 5^4 , 5^6 -tetra(methylsulfonyloxy)-2,4,6,8-tetramethyl-1,3,5,7(1,3)tetrabenzenacyclooctaphane (2l). Resorcinarene 2j (1.4 g, 1 mmol) was dissolved in a 1:1 solvent mixture of THF and ethanol (60 mL). 10% Palladium on carbon (100 mg, 0.1 mmol) was added and the solution stirred at room temperature under a positive pressure of hydrogen. After 6 h the reaction was checked for completion (tlc) and the catalyst filtered off through Celite®, washing with dichloromethane (100 mL). Evaporation of the solvent afforded tetramesylate 2l (850 mg, 99%) as an off-white solid. Mp $>$ 300 °C (dec.); $v_{\text{max}}/\text{cm}^{-1}$ (KBr) 3494s + 3389s (O–H, H-bonded), 3032s (C–H, aromatic), $2968s + 2933s$ (CH₃), $1619m + 1520s$ (aryl stretch), $1354s + 1184s$ (–SO₂–) $\delta_{\rm H}$ (300 MHz, DMSO- d_6) 1.42 (12H, d, J=6.9 Hz, –CH₃), 2.93 (12H, s, $-SO_2CH_3$), 4.63 (4H, q, J = 6.9 Hz, H-2,4,6,8), 6.12 (2H, s, H_1^2 , $\tilde{5}^2$), 6.42 (2H, s, 1^5 , 5^5), 7.18 (2H, s, 3^5 , 7^5), 7.48 (2H, s, $(3^2,7^2)$, 9.16 (4H, s, –OH); δ_C (75 MHz, DMSO- d_6) 20.3 (CH_3) , 29.7 (C-2,4,6,8), 37.0 (SO₂CH₃), 102.0 (C-1⁵,5⁵), 114.0 $(C-3^5,7^5)$, 122.6 $(C-1^1,1^3,5^1,5^3)$, 125.2 $(C-1^2,5^2)$, 126.8 $(C-3^2,7^2)$, 136.3 $(C-3^1,3^3,7^1,7^3)$, 144.4 (7^2) , 136.3 $(C-3^1,3^3,7^1,7^3)$ $(C-3^4,3^6,7^4,7^6)$, 152.4 $(C-1^4,1^6,5^4,5^6)$; MALDI-TOF m/z (rel. int.) 896.04 $[M^+ + K^+]$ (48), 879.94 $[M^+ + Na^+]$ (100). Found: C, 50.28; H, 4.64; S, 14.85%; C₃₆H₄₀O₁₆S₄ requires C, 50.46; H, 4.70; S, 14.97%.

6.2. General procedure for the synthesis of bridged resorcinarenes (3)

Tetraprotectedresorcinarene 2 (0.25–0.5 mmol, 1 equiv), diamine 1, 6 or 12 (1.2 equiv) and paraformaldehyde (10 equiv) were added to a high-pressure glass reaction

vessel with a screw top. Acetonitrile (40 mL) was added and the vessel sealed and placed in an oil bath at 90° C. After 40 min the reaction was checked by tlc for complete consumption of 2. Silica gel $(1-2 \rho)$ was then added and the solvent removed under reduced pressure. The solidsupported crude material was then finely ground and subjected to column chromatography (30 g silica gel eluting with ethyl acetate–petroleum ether 1:9–1:1) to furnish bridged derivatives 3.

6.2.1. $(R,R)-1^2,1^4,11^2,11^4$ -Tetrahydroxy-12,14,15,17tetra-methyl-3,9-bis(1-phenylethyl)-13⁴,13⁶,16⁴,16⁶tetra(2',4',6'-triisopropylbenzenesulfonyloxy)-3,9-diaza-1,11(1,3,5),13,16(1,3)tetrabenzenabicyclo[9.3.3]heptadecaphane (3f). Yield 78%. Mp $232-233$ °C (from diethyl ether–petroleum ether); $\lbrack \alpha \rbrack_{D}$ +67.6 (c 1.40 in CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3516s (O–H, H-bonded), 2964s+ $2872s$ (C-H), $1599m+1481m$ (aryl stretch), $1376s+$ 1180s (-SO₂-O-); δ_H (300 MHz, CDCl₃) 0.99 (2H, m, H-6), 1.11 (4H, m, H-5,7), 1.18 (24H, d, $J=6.6$ Hz, $7'/9'$ *i*-Pr), 1.31 (60H, br d, $J=6.6$ Hz, $7'/8'/9'$ *i*-Pr + -CH₃), 1.37 (6H, d, $J=6.6$ Hz, NCHC H_3), 2.12 and 2.35 (4H, m, H-4,8), 2.96 $(4H, m, H-8')$, 3.66 and 3.77 (4H, d, $J_{AB} = 15.4$ Hz, H-2,10), 3.74 (2H, m, NCHCH₃), 4.13 (8H, m, H-7 $\frac{1}{9}$), 4.52 and 4.54 (4H, $2 \times q$, $J=6.6$ Hz, H-12,14,15,17), 6.74 (2H, s, $H-13⁵,16⁵$), 6.80 (2H, s, H-13²,16²), 7.09 (2H, s, H-1⁶,11⁶), 7.18–7.31 (18H, m, Ph + H-3',5'); δ_C (75 MHz, CDCl₃) 18.6 $(NCHCH₃), 21.6$ and 22.3 (– $CH₃$), 23.1 (C-6), 23.5 (8^t i-Pr), 24.7 $(7'/9'$ *i*-Pr), 26.3 (C-5,7), 30.0 (C-7 $'/9'$), 31.0 and 31.3 $(C-12, 14, 15, 17)$, 34.3 $(C-8)$, 48.4 $(C-2, 10)$, 49.5 $(C-4, 8)$, 61.7 (NCHCH₃), 108.4 (C-1³,11³), 115.6 (C-13⁵,16⁵), 118.1 and 119.1 $(C^{-1}, 1^5, 11^1, 11^5)$, 123.7 $(C^{-1}, 11^6)$, 124.0 and 124.1 (C-3['],5[']), 127.3 (C-13²,16²), 128.0 and 128.5 (Ph), 130.3 and 130.6 $(C-1')$, 139.6 and 140.2 $(C-13¹, 13³)$, $16¹,16³$), 142.2 (Ph), 144.5 and 144.6 (C-13⁴,13⁶,16⁴,16⁶), 151.0 and 151.2 $(C-2^{\prime},6^{\prime})$, 153.0 $(C-1^2,1^4,11^2,11^4)$, 154.3 and 154.4 (C-4'); MALDI TOF: m/z (rel. int.) 1942.89 [M⁺] (53), 1838.6 $[M^+ - PhCHCH_3]$ (10). Found: C, 71.04; H, 7.72; N, 1.46; S, 6.38%; C₁₁₅H₁₅₀N₂O₁₆S₄ requires C, 71.03; H, 7.77; N, 1.44; S, 6.60%

6.2.2. $(R,R)-1^2,1^4,11^2,11^4$ -Tetrahydroxy-13⁴,13⁶,16⁴,16⁶tetra(4-methylbenzoyloxy)-12,14,15,17-tetramethyl-3,9 bis(1-phenylethyl)-3,9-diaza-1,11(1,3,5),13,16(1,3)tetrabenzenabicyclo[9.3.3]heptadecaphane (3g). Yield 55%. Mp 193–195 \degree C (from diisopropyl ether–dichloromethane); $\left[\alpha\right]_{\text{D}}$ +7.8 (c 2.64 in CHCl₃); $\nu_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3480s (O–H, H-bonded), 2972s + 2873s (C–H, aliphatic), 1729s (C=O, ester); δ_H (300 MHz, CDCl₃) 0.93 (2H, m, H-6), 1.10 (4H, m, H-5,7), 1.40 (6H, d, $J=6.7$ Hz, NCHCH₃), 1.54 (12H, d, $J=6.7$ Hz, $-CH_3$), 2.19 and 2.33 (4H, m, H-4,8), 2.46 (12H, s, H-7[']), 3.78 (6H, br, NCHCH₃ and H-2,10), 4.49 (4H, br m, H-12, 14, 15, 17), 7.04 (2H, s, H-13², 16^2), 7.14 (2H, s, H-13⁵,16⁵), 7.22–7.30 (12H, br m, $H-1^6$,11⁶ + Ph) 7.34 (8H, m, $H-3^7$, 5'), 8.23 (8H, m, $H-2^7$, 6'); δ_C (100 Hz, CDCl₃) 17.7 (NCHCH₃), 21.7 (C-7[']), 22.2 and 22.4 ($-CH_3$), 24.8 (C-5,7), 25.9 (C-6), 30.6 + 30.7 (C-12,14,15,17), 49.8 (C-2,10), 52.0 (C-4,8), 62.3 (NCHCH₃), 108.8 (C-1³,11³), 115.8 (C-13⁵,16⁵), 119.8 $(C-1^1, 1^5, 1^1^1, 1^1^5), 122.7 (C-1^6, 11^6), 126.9 (C-1^7), 127.0$ (Ph), 127.2 (C-13²,16²), 128.0 + 128.4 (Ph), 129.4 (C-3',5'), 130.5 (C-2^{*'*},6[']), 137.7 (C-13¹,13³,16¹,16³), 141.9 (Ph), 144.6 (C-4'), 146.1 (C-13⁴,13⁶,16⁴,16⁶), 152.9 (C-1²,1⁴,

 $11^2,11^4$), 165.8 (C=O); MALDI TOF: m/z (rel. int.) 1350.65 $[M^+]$ (79), 1247.25 (44). Found: C, 77.53; H, 6.54; N, 2.03%; $C_{87}H_{86}N_2O_{12}$ requires C, 77.31; H, 6.41; N, 2.07%.

6.2.3. (R, R) -13⁴,13⁶,16⁴,16⁶-Tetra(benzyloxycarbonyloxy)-1²,1⁴,11²,11⁴-tetrahydroxy-12,14,15,17-tetramethyl-3,9bis(1-phenylethyl)-3,9-diaza-1,11(1,3,5),13,16(1,3)tetrabenzenabicyclo[9.3.3]heptadecaphane (3h). Yield 70%. Mp 117–119 °C (from diethyl ether); $[\alpha]_D$ +6.7 (c 1.00, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3502s (O–H, H-bonded), 2971s (C–H, aliphatic), 1757s (C=O, ester); $\delta_{\rm H}$ (300 MHz, CDCl₃) 0.77 $(2\overline{H},$ qu, $J=7.1$ Hz, H-6), 0.98 (4H, qu, $J=7.1$ Hz, H-5,7), 1.40 (6H, d, $J=6.5$ Hz, NCHCH₃), 1.45 (12H, d, $J=6.5$ Hz, $-CH_3$), 2.06 and 2.23 (4H, m, H-4,8), 3.71 (4H, s, H-2,10), 3.79 (2H, q, $J=6.5$ Hz, H- α), 4.43 (4H, m, H-12,14,15,17), 5.32 and 5.34 (8H, 2 \times s, H-2'), 6.85 (2H, s, H-13²,16²), 7.04 $(2H, s, H-13⁵, 16⁵), 7.19 (2H, s, H-1⁶, 11⁶), 7.20–7.48 (30H, m,$ Ph); δ_C (75 Hz, CDCl₃) 18.0 (NCHCH₃), 22.0 and 22.1 (–CH3), 24.8 (C-5,7), 25.5 (C-6), 30.5 (C-12,14,15,17), 49.5 $(C-2,10)$, 51.6 $(C-4,8)$, 62.1 (NCHCH₃), 70.6 (2'), 109.0 $(C-1^3, 1^3)$, 115.2 $(C-13^5, 16^5)$, 119.5 $(C-1^1, 1^5, 11^1, 11^5)$, 123.0 $(C-1^6, 11^6)$, 127.2 $(C-13^2, 16^2)$, 127.4 + 128.1 + 128.4 + $128.5 + 128.7$ (Ph), 135.0 (Ph), 137.7 (C-13¹,13³,16¹,16³), 141.5 (Ph), 146.3 (C-13⁴,13⁶,16⁴,16⁶), 152.9 (C-1²,1⁴) $11^2,11^4$), 154.1 (C-1'); MALDI-TOF m/z (rel. int.) 1416.36 $[M^+ + 2H]$ (100). Found: C, 73.88; H, 6.17; N, 2.02%; $C_{87}H_{86}N_2O_{16}$ requires C, 73.82; H, 6.12; N, 1.98%.

6.2.4. (R,R) -13⁴,13⁶,16⁴,16⁶-Tetraacetoxy-1²,1⁴,11²,11⁴tetra-hydroxy-12,14,15,17-tetramethyl-3,9-bis(1-phenylethyl)-3,9-diaza-1,11(1,3,5),13,16(1,3)tetra-benzenabicyclo- [9.3.3]heptadecaphane (3k). Yield 20%. Mp 162-164 °C (from diethyl ether - petroleum ether); $[\alpha]_D$ +8.1 (c 1.62, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3502s (O–H, H-bonded), $2972s + 2874s$ (C–H, aliphatic), 1747s (C=O, ester); δ_H (300 MHz, CDCl3) 0.84 (2H, m, H-6), 0.97 (4H, m, H-5,7), 1.40 (6H, d, $J=7.0$ Hz, NCHCH₃), 1.48 (12H, d, $J=7.1$ Hz, $-CH_3$), 2.08 and 2.24 (4H, m, H-4,8), 2.39 (12H, s, OC(O)CH₃), 3.73 (4H, s, H-2,10), 3.81 (2H, q, $J=7.0$ Hz, H-a), $4.35(4H, m, H-12, 14, 15, 17)$, 6.85 (2H, s, H-13⁵, 16⁵), 6.89 (2H, s, H-13²,16²), 7.28 (2H, s, H-1⁶,11⁶), 7.19–7.34 (10H, m, Ph); δ_c (75 MHz, CDCl₃) 17.5 (NCHCH₃), 21.0 $(OC(O)CH₃)$, 22.0 and 22.1 (–CH₃), 25.0 and 25.7 (C-5, 6,7), 30.5 (C-12,14,15,17), 49.4 (C-2,10), 52.0 (C-4,8), 62.2 $(NCHCH₃), 109.2 (C-1³,11³), 115.9 (C-13⁵,16⁵), 119.9$ $(C-1^1, 1^5, 1^1^1, 11^5)$, 122.7 $(C-1^6, 11^6)$, 127.1 (Ph), 127.3 $(C-13^2, 16^2)$, 128.0 and 128.4 (Ph), 137.1 and 137.2 $(C-13¹, 13³, 16¹, 16³), 141.5$ (Ph), 145.8 and 145.9 (C-13⁴, 13⁶,16⁴,16⁶), 153.0 (C-1²,1⁴,11²,11⁴), 170.0 (C=O); MALDI-TOF m/z (rel. int.) 1047.75 $[M^+]$ (100), 1003.60 $[M^+ - COCH_3]$ (20), 943.60 $[M^+ - CHCH_3Ph]$ (75). Found: C, 72.17; H, 6.59; N, 2.75%; $C_{63}H_{70}N_2O_{12}$ requires C, 72.26; H, 6.74; N, 2.67%.

6.2.5. (R,R) -1²,1⁴,11²,11⁴-Tetrahydroxy-12,14,15,17-tetramethyl-13⁴ ,13⁶ ,16⁴ ,16⁶ -tetra(methylsulfonyl-oxy)-3, 9-bis(1-phenylethyl)-3,9-diaza-1,11(1,3,5),13,16(1,3) tetra-benzenabicyclo[9.3.3]hepta-decaphane (3l). Yield 51%. Mp 201-203 °C (from dichloromethane-diethyl ether); $[\alpha]_D$ +8.1 (c 2.13, CHCl₃); $\nu_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3523s (O-H, H-bonded), $2975s + 2875s$ (C-H), $1606m +$ 1481m (aryl stretch), $1370s+1183s$ (–SO₂–O–); δ_{H}

 $(300 \text{ MHz}, \text{CDCl}_3)$ 1.05 (2H, m, H-6), 1.19 (4H, m, H-5,7), 1.37 (6H, d, $J=6.8$ Hz, NCHCH₃), 1.50 (12H, d, $J=7.1$ Hz, $-CH_3$), 2.18 and 2.37 (4H, m, H-4,8), 2.37 and 2.40 (12H, $2 \times s$, –SO₂Me), 3.78 (4H, s, H-2,10), 3.81 (2H, q, J= 6.8 Hz, NCHCH₃), 4.56 and 4.59 (4H, $2 \times q$, $J = 7.1$ Hz, H-12,14, 15,17), 6.80 (2H, s, H-13²,16²), 7.15 (2H, s, H- 1^6 , 11^6), $7.17 - 7.32$ (10H, m, Ph), 7.47 (2H, s, H- 13^5 , 16^5); δ_c $(75 \text{ MHz}, \text{CDCl}_3)$ 16.8 (NCHCH₃), 21.8 and 22.0 (-CH₃), 24.5 (C-5,7), 25.0 (C-6), 31.7 (C-12,14,15,17), 38.1 $(SO₂Me)$, 48.9 (C-2,10), 49.8 (C-4,8), 61.1 (NCHCH₃), 108.5 (C-1³,11³), 116.4 (C-13⁵,16⁵), 118.2 (C-1¹,1⁵) $11¹,11⁵$), 124.3 (C-1⁶,11⁶), 127.1 (C-13²,16²), 127.5 127.9 and 128.4 (Ph), 140.2 (C- $13¹, 13³, 16¹, 16³$), 141.3 (Ph), 144.6 $(C-13^4, 13^6, 16^4, 16^6)$, 153.6 $(C-1^2, 1^4, 11^2, 11^4)$; MALDI-TOF m/z (rel. int.) 1192.1 [M⁺ + 2H] (100). Found: C, 59.48; H, 5.98; N, 2.36; S, 10.52%; $C_{59}H_{70}N_2O_{16}S_4$ requires C, 59.48; H, 5.92; N, 2.35; S, 10.76%.

6.2.6. Heptanedioic acid bis[(1-phenylethyl)amide] (5). Pimelic acid 4 (641 mg, 4 mmol) and hydroxybenzotriazole (270 mg, 2 mmol) were dissolved in dry dichloromethane (50 mL) and cooled to 0 °C. Dicyclohexylcarbodiimide (2.06 g, 10 mmol) was added, followed by slow addition of (R) - α -methylbenzylamine (1.08 mL, 8.4 mmol). The reaction was allowed to warm to room temperature slowly overnight. On completion, the reaction was added to aqueous sodium carbonate (100 mL) and extracted with dichloromethane $(3 \times 70 \text{ mL})$. The organic layers were combined and dried with magnesium sulphate and then reduced to yield a crude white product, which was purified by column chromatography (60 g silica gel, eluting with methanol–ethyl acetate 2:98), to afford diamide 5 (1.40 g, 95%) as a colourless powder. Mp $157-158$ °C (ethyl acetate–ethanol); $[\alpha]_D$ + 108.5 (c 2.50, EtOH); $\nu_{\text{max}}/\text{cm}^{-1}$ (CH_2Cl_2) 1670s (C=O stretch, amide I), 1506s (C=O, amide II), 1262s; δ_H (300 MHz, CDCl₃) 1.33 (2H, m, H-4), 1.46 (6H, d, $J=6.9$ Hz, $-CH_3$), 1.62 (4H, m, H-3,5), 2.14 $(4H, m, H-2, 6), 5.10 (2H, q, J=6.9 Hz, H-\alpha), 5.86 (2H, br d,$ $J = \sim$ 7 Hz, NH), 7.21–7.35 (10H, m, Ph); δ_C (75 MHz, CDCl3) 21.8 (–CH3), 25.1 (C-3,5), 28.5 (C-4), 36.3 (C-2,6), 48.6 (C-a), 126.1, 127.3, 128.6 and 143.4 (Ph), 171.9 $(C=0)$; m/z (rel. int.) 367.2 [M⁺ + H] (57), 263.2 (8), 246.2 (9), 142 (12), 120 [PhCH(NH)CH3] (18), 105 [PhCHCH3] (100) , 91 [PhCH₂] (47). Found C, 75.49; H, 8.26; N, 7.60%; $C_{23}H_{30}N_2O_2$ requires C, 75.38; H, 8.25; N, 7.64%.

6.2.7. N, N' -Bis(1-phenylethyl)heptane-1,5-diamine (6). Di-amide 5 (4.30 g, 11.7 mmol) was dissolved in dry THF (150 mL) and cooled to 0° C. Lithium aluminium hydride (1.78 g, 46.9 mmol) was added portion-wise to the reaction, which was then heated to reflux for 18 h. After the reaction was complete (tlc), it was cooled to 0° C and quenched by the slow addition of water (1.7 mL), 1 M sodium hydroxide (3.4 mL) followed by water (3.4 mL). The reaction was stirred at room temperature for 1 h, the salts removed via filtration through Celite[®], washing with dichloromethane (100 mL) , to afford diamine 6 $(3.78 \text{ g}, 95\%)$ as a clear oil, which was judged to be $>95\%$ pure for the next step. The product was characterized as its ditosylate. Mp $137-138$ °C (ethanol–dichloromethane); $[\alpha]_D$ +18.8 (c 2.6, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3024s (C=C–H), 2940s + 2856s (C–H, aliphatic), 1599s (aryl), $1380s+1162s$ (SO₂–O); $\delta_{\rm H}$ $(400 \text{ MHz}, \text{ CDCl}_3)$ 0.84 (6H, br m, H-3,4,5), 1.06 and

1.28 (4H, m \times 2, H-2,6), 1.35 (6H, d, J=7.1 Hz, –CH₃ \times 2), 2.41 (6H, s, -Tos-C $H_3 \times 2$), 2.92 (4H, m, H-1,7), 5.15 (2H, q, $J=7.1$ Hz, H- α), $7.18-7.25$ (10H, m, Ph), 7.28 (4H, d, \bar{J} =8.0 Hz, –Tos), 7.71 (4H, d, J =8.0 Hz, -Tos); δ_C $(75 \text{ MHz}, \text{ CDCl}_3)$ 16.6 $(-CH_3)$, 21.4 $(Tos-CH_3)$, 26.6 $(C-3,5)$, 28.2 $(C-4)$, 30.4 $(C-2,6)$, 44.1 $(C-1,7)$, 55.2 $(C-\alpha)$, 127.1, 127.5, 128.2, 129.5, 138.4, 140.3 and 142.9 (Ph); m/z (rel. int.) 669 $[M^+ + Na]$ (100). Found C, 68.68; H, 7.16; N, 4.32; S, 9.92%; C₃₇H₄₆N₂O₄S₂ requires C, 68.70; H, 7.17; N, 4.33; S, 9.91%.

6.2.8. 4-Oxo-heptanedioic acid bis[(1-phenylethyl) amide] (10). Oxo-pimelic acid 9 (3.49 g, 20 mmol) and hydroxybenzotriazole (1.35 g, 10 mmol) were dissolved in dry dichloromethane (200 mL) and cooled to 0° C. Dicyclohexylcarbodiimide (10.3 g, 50 mmol) was added, followed by slow addition of (R) - α -methylbenzylamine (5.4 mL, 42 mmol). The reaction was allowed to warm to room temperature slowly overnight. On completion, the reaction was added to aqueous sodium carbonate (200 mL) and extracted with dichloromethane $(3 \times 100 \text{ mL})$. The organic layers were combined and dried with magnesium sulphate and then reduced to yield a crude white product, which was purified by column chromatography (100 g silica gel, eluting with methanol–ethyl acetate 2:98). In this way, diamide 10 (6.30 g, 83%) was obtained as a white powder. Mp 160–161 °C (ethyl acetate–ethanol); $[\alpha]_D$ + 121.5 $(c$ 2.20, EtOH); v_{max}/cm^{-1} (CHCl₃) 3432m (N–H), 1716s (C=O, ketone), 1672s (C=O, amide), 1508s (NH bend); $\delta_{\rm H}$ $(400 \text{ MHz}, \text{CDCl}_3)$ 1.45 (6H, d, $J=6.9 \text{ Hz}, 2 \times \text{NCHCH}_3$), 2.45 (4H, m, H-3,5), 2.76 (4H, m, H-2,6), 5.06 (2H, q, $J=$ 7.0 Hz, $2 \times H-\alpha$), 5.94 (2H, br d, $J=6.9$ Hz, $2 \times N-H$), 7.21–7.34 (10H, m, Ph); δ_C (100 MHz, CDCl₃) 21.8 (-CH₃), 30.2 (C-3,5), 37.6 (C-2,6), 48.9 (C-a), 126.1, 127.2, 128.6 and 143.3 (Ph), 170.7 (C-1,7), 208.8 (C-4); m/z (rel. int.) 381.2 $[M^+ + H]$ (42), 260.1 $[M^+ - PhCH(CH_3)NH]$ (17), 156.0 (21), 120.0 [PhCH(CH3)NH] (25), 105.0 [PhCHCH3] (100). Found C, 72.58; H, 7.26; N, 7.29%; C₂₃H₂₈N₂O₃ requires C, 72.61; H, 7.42; N, 7.36%.

6.2.9. 4,4-Dimethoxy-heptanedioic acid bis[(1-phenylethyl)-amide] (11). Diamide 10 (0.98 g, 2.5 mmol) was dissolved in dry methanol (30 mL) with a catalytic amount of p-toluenesulfonic acid (40 mg, 0.2 mmol). Trimethyl orthoformate (2 mL, 18 mmol) was added and the reaction heated to gentle reflux overnight. On completion of the reaction, solid sodium carbonate was added and the methanol reduced on a rotary evaporator. Ethyl acetate (100 mL) was added to the crude material, which was then washed with water (25 mL) and brine (25 mL). The organic layer was then dried with magnesium sulphate and reduced to give a crude oil which was purified via column chromatography (50 g silica gel, eluting with methanol– ethylacetate 2:98), to give dimethoxyketal diamide 11 (0.98 g, 92%) as fine white needles. Mp 129–131 °C (ethyl acetate–ethanol); $[\alpha]_D$ +86.9 (c 2.05, EtOH); $\nu_{\text{max}}/\text{cm}^-$ (CHCl₃), 1663s (C=O, amide), 1505s (N–H bend); δ_H $(400 \text{ MHz}, \text{CDCl}_3)$ 1.45 (6H, d, $J=6.9 \text{ Hz}, 2 \times \text{NCHCH}_3$), 1.89 (4H, m, H-3,5), 2.19 (4H, m, H-2,6), 3.13 (6H, s, OCH₃), 5.03 (2H, q, $J=7.0$ Hz, $2\times$ H- α), 6.13 (2H, br d, $J=7.7$ Hz, $2\times$ N–H), 7.18–7.33 (10H, m, Ph); δ_C (100 MHz, CDCl3) 21.8 (–CH3), 28.1 (C-3,5), 31.2 $(C-2,6)$, 47.9 $(-OCH_3)$, 48.8 $(C-\alpha)$, 102.5 $(C-4)$, 126.1,

127.3, 128.6 and 143.2 (Ph), 171.7 (C-1,7); m/z (rel. int.) 449.2 $[M^+ + Na]$ (5), 395.2 $[M^+ - OCH_3]$ (56), 259.1 (19), 142.0 (15), 105.0 [PhCHCH3] (100). Found C, 70.71; H, 8.01; N, 6.74% ; C₂₅H₃₄N₂O₄ requires C, 70.40; H, 8.03; N, 6.57%.

6.2.10. $4,4$ -Dimethoxy- N,N' -bis-(1-phenylethyl)heptane-1,7-diamine (12). Diamide 11 (1.11 g, 2.6 mmol) and triethylamine (1.8 mL, 13 mmol) were added to dry diethyl ether (100 mL) and cooled to 0° C. Lithium aluminium hydride (395 mg, 10.4 mmol) was added to the solution in small portions. Thereafter, the reaction was heated to reflux for 4–6 h until reduction was complete (tlc control). The reaction was then cooled to 0° C and quenched by the addition of H_2O (0.4 mL), 1 M NaOH (0.8 mL) and H_2O (0.8 mL) in that order. The heterogeneous solution was allowed to stir at room temperature for approximately one hour until all the salts had turned white. These were then filtered off through Celite[®] and washed with dichloromethane (200 mL). The filtrate was then reduced to afford diamine 12 (880 mg, 85%) as an oil. $[\alpha]_D$ +37.5 (c 2.92, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3013s (aromatic C–H), $2954s+2831s$ (aliphatic C–H), 1602w (aryl ring), 1188m (C–N), 1113s (C–O–C); δ_H (400 MHz, CDCl₃) 1.39 (3H, d, $J=6.6$ Hz, $-CH_3$), 1.32–1.61 (10H, m, H-2,3,5,6 + NH \times 2), $2.42 + 2.50$ (4H, $2 \times dt$, $J = 11.4$, 7.1 Hz, H-1,7), 3.08 (6H, s, OCH₃), 3.78 (2H, q, $J=6.6$ Hz, NCHCH₃), 7.20–7.35 (10H, m, Ph); δ_C (100 MHz, CDCl₃) 23.9 (CH₃), 24.1 (C-2,6), 30.2 (C-3,5), 47.6 (C-1,7+OMe), 58.4 (NCHCH₃), 103.1 (C-3), 126.6, 127.0, 128.4 and 144.8 (Ph).

6.2.11. (R,R) -1²,1⁴,13²,13⁴-Tetrahydroxy-14,16,17,19tetra-methyl-3,11-bis(1-phenylethyl)-15⁴,15⁶,18⁴,18⁶-tetra-(p-toluenesulfonyloxy)-3,11-diaza-1,13(1,3,5),15,18(1,3) tetrabenzenabicyclo[11.3.3]nona-decaphane(13a).

Tetratosylresorcinarene 2a (465 mg, 0.5 mmol), diamine 6 (203 mg, 0.6 mmol) and paraformaldehyde (300 mg, 10 mmol) were added to a high-pressure glass reaction vessel with a screw top. Acetonitrile (40 mL) was added and the vessel sealed and placed in an oil bath at 80° C. After two hours the reaction was checked by tlc for complete consumption of $2a$. Silica gel $(2-3 g)$ was then added and the solvent removed under reduced pressure. The solidsupported crude material was then finely ground and subjected to column chromatography (40 g silica gel eluting with ethyl acetate–petroleum ether 3:7). In this manner, bridged resorcinarene 13a (220 mg, 36%) eluted first, followed by dimer 14 (80 mg, 5%). Mp $133-137$ °C (diisopropyl ether–dichloromethane); $\lbrack \alpha \rbrack_{D} + 13.2$ (c 2.40, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃): 3528s (O–H, H-bonded), $2974m+2932m$ (C–H, aliphatic), 1598s (aryl-H), $1371s+$ 1176s (-SO₂-O-); δ_H (300 MHz, CDCl₃) 1.07-1.44 (10H, m, H-5,6,7,8,9), 1.30 (6H, d, $J=6.9$ Hz, NCHCH₃), 1.41 (12H, d, $J=7.0$ Hz, $-CH_3$), 1.99 and 2.40 (4H, m, H-4,10), 2.47 and 2.49 (12H, $2 \times s$, H-7[']), 3.60 and 3.69 (4H, AB d, J_{AB} = 14.6 Hz, H-2,12), 3.79 (2H, q, J = 6.9 Hz, H- α), 4.52 and 4.55 (4H, $2 \times q$, $J = 7.0$ Hz, H-14,16,17,19), 6.45 (2H, s, H-15²,18²), 6.78 (2H, s, H-15⁵,18⁵), 7.07 (2H, s, H-1⁶,13⁶), 7.17–7.33 (10H, m, Ph), 7.39 and 7.41 (8H, $2 \times d$, $J=$ ~8.0 Hz, H-3',5'), 7.94 (8H, 2×d, J = ~8.0 Hz, H-2',6'); δ_C (75 MHz, CDCl₃) 14.1 (NCHCH₃), 20.6 and 20.7 $(-CH₃), 21.7$ $(C-7')$, 25.0, 25.8 and 26.4 $(C-5,6,7,8,9)$, 31.8 and 31.9 (C-14,16,17,19), 46.4 (C-2,12), 48.4 (C-4,10),

57.1 (C- α), 108.8 (C- 1^3 , 13³), 114.8 (C- 15^5 , 18⁵), 118.5 and 118.9 (C- $1^1, 1^5, 13^1, 13^5$), 123.6 (C- $1^6, 13^6$), 127.0 (Ph), 127.4 $(C-15^2, 18^2)$, 128.0 (Ph), 128.4, 128.5 and 128.6 $(C-2^{\prime}, 6^{\prime} +$ Ph), 130.1 (C-3',5'), 132.9 (C-1'), 139.8 and 140.0 (C-15¹, 15^3 , 18^1 , 18^3), 140.2 (Ph), 144.8 and 145.0 (C- 15^4 , 15^6 , 18^4 , 18^6), 145.6 (C-4'), 153.1 and 153.8 (C- 1^2 , 1^4 , 13^2 , 13^4); MS: m/z (rel. int.) 1523.4 [M⁺] (55), 1417.3 [M⁺ – H–PhCHCH3] (13), 1370.3 (7), 1185.1 (7), 1028.3 (3), 875.1 (3), 547.1 (7), 339.3 (57), 105.1 [PhCHCH₃] (100). Found C, 66.98; H, 5.98; N, 1.84; S, 8.05%; C₈₅H₉₀N₂O₁₆S₄ requires C, 66.99; H, 5.95; N, 1.84; S, 8.42%.

6.2.12. (R, R, R, R) -1²,1⁴,13²,13⁴,17²,17⁴,29²,29⁴-Octahydroxy-14,16,30,32,33,35,36,38-octamethyl-3,11,19,27 tetra(1-phenylethyl)-15⁴,15⁶,31⁴,31⁶,34⁴,34⁶,37⁴,37⁶-octa-(p-toluenesulfonyloxy)-3,11,19,27-tetraaza-1,13,17,29 $(1,3,5),15,31,34,37(1,3)$ tricyclo $[27.3.3.3^{13,17}]$ octatriaconta-phane (14). From synthesis of bridged resorcinarene 13a. Mp $161-165$ °C (diisopropyl ether-dichloromethane); $[\alpha]_D$ –7.0 (c 1.33 in CHCl₃); $\nu_{\text{max}}/\text{cm}^{-1}$ (CHCl₃): 3527s (O–H, H-bonded), 3022s (C–H, aromatic), 2930m (C–H, aliphatic), 1598s (aryl stretch), $1371s+1177s$ (–SO₂–); $\delta_{\rm H}$ $(300 \text{ MHz}, \text{CDCl}_3)$ 1.03 (4H, m, H-7,23), 1.21 (12H, d, J= 6.8 Hz, NCHC H_3), 1.27 (8H, m, C-5,6,8,9,21,22,24,25), 1.36 and 1.39 (24H, d, $J=6.7$ Hz, $-CH_3$), 2.02 and 2.40 (8H, br m, H-4,10,20,26), 2.45 (24H, s, H-7[']), 3.34 (8H, s, H-2,12,18,28), 3.65 (4H, q, $J=6.8$ Hz, NCHCH₃), 4.43 and 4.48 (8H, $2 \times q$, $J=7.0$ Hz, H-14,16,30,32,33,35,36,38), 6.44 (4H, s, H-15²,31²,34²,37²), 6.72 (4H, s, H-15⁵,31⁵, $34^5,37^5$, 7.00 (4H, s, H-1⁶,13⁶,17⁶,29⁶), 7.02–7.17 (20H, m, Ph), 7.35 (16H, d, $J = \sim 8.1$ Hz, H-3',5'), 7.88 (16H, d, $J =$ ~8.1 Hz, H-2',6'); δ_C (75 MHz, CDCl₃) 16.0 (NCHCH₃), $20.4 + 20.6$ (-CH₃), 21.7 (C-7[']), 26.7, 27.4 and 30.0 (C-5,6,8,9,21,22,24,25), 31.7 and 31.9 (C-14,16,30,32,33, 35, 36,38), 46.9 (C-2,12,18,28), 49.3 (C-4,10,20,26), 58.1 $(C=NCHCH₃), 109.0 (C-1³,13³,17³,29³), 114.5 (C-15⁵,31⁵$ 34^5 , 37⁵), 118.9 (C-1¹, 1⁵, 13¹, 13⁵, 17¹, 17⁵, 29¹, 29⁵), 123.3 $(C-1⁶, 13⁶, 17⁶, 29⁶)$, 127.2 (Ph), 127.4 ($C-15², 31², 34², 37²)$, 128.1 (Ph), 128.4, 128.4 and 128.5 $(C-2', 6' + Ph)$, 130.0 $(C-3',5')$, 133.2 and 133.4 $(C-1')$, 139.7 and 140.1 $(C-15^1, 15^3, 31^1, 31^3, 34^1, 34^3, 37^1, 37^3)$, 140.4 (Ph), 144.9 and 145.1 $(C-15^4, 15^6, 31^4, 31^6, 34^4, 34^6, 37^4, 37^6)$, 145.5 and 145.6 (C-4'), 153.5 and 153.8 (C-1², 1⁴, 13², 13⁴, 17², $17^4,29^2,29^4$); MALDI TOF: m/z (rel. int.) 3047.3 [M⁺] (15), 1524.7 $[M^+/2+H]$ (72). Found C, 66.52; H, 5.96; N, 1.86; S, 8.16%; $C_{170}H_{180}N_4O_{32}S_8 \cdot H_2O$ requires C, 66.60; H, 5.98; N, 1.83; S, 8.37%.

6.2.13. (R,R) -1²,1⁴,13²,13⁴-Tetrahydroxy-14,16,17,19tetra-methyl-7,7-dimethoxy-3,11-bis(1-phenyl-ethyl)- 15⁴,15⁶, 18⁴,18⁶-tetra(p-toluenesulfonyloxy)-3,11-diaza-1,13(1,3,5),15,18(1,3)tetrabenzenabicyclo[11.3.3]nonadecaphane (13b). Tetratosylresorcinarene 2a (580 mg, 0.5 mmol), diamine 12 (590 mg, 1.48 mmol) and paraformaldehyde (300 mg, 10 mmol) were added to a highpressure glass reaction vessel with a screw top. Acetonitrile (40 mL) was added and the vessel sealed and placed in an oil bath at 80 \degree C. After 1 1/2 h the reaction was checked by tlc for complete consumption of $2a$. Silica gel $(2-3 g)$ was then added and the solvent removed under reduced pressure. The solid-supported crude material was then finely ground and subjected to column chromatography (50 g silica gel eluting with ethyl acetate–petroleum ether 3:7), to afford

bridged resorcinarene 13b (450 mg, 57%). Mp 146–148 °C (diisopropyl ether–dichloromethane); $\lbrack \alpha \rbrack_{D} + 1.0$ (c 2.40, CHCl₃); $v_{\text{max}}/\text{cm}^{-1}$ (CHCl₃): 3526s (O–H, H-bonded), $2971m$ (C-H, aliphatic), 1599s (aryl stretch), $1371s+1177s$ $(-SO₂–)$, 1092s (C–O–C, ether); δ_H (300 MHz, CDCl₃) 1.29 (6H, d, $J=6.9$ Hz, NCHC H_3), 1.26–1.37 (8H, m, H-5,6, 8,9), 1.42 (12H, d, $J=7.1$ Hz, $-CH_3$), 2.02 and 2.45 (4H, m, H-4,10), 2.48 and 2.49 (12H, $2 \times s$, H-7'), 3.02 (6H, s, $-OCH_3$), 3.60 and 3.66 (4H, AB d, J_{AB} = 14.8 Hz, H-2,12), 3.80 (2H, q, $J=6.9$ Hz, H- α), 4.52 and 4.57 (4H, $2 \times q$, $J=$ 7.1 Hz, H-14,16,17,19), 6.44 (2H, s, H-15²,18²), 6.73 (2H, s, $H-15⁵, 18⁵$), 7.07 (2H, s, $H-1⁶, 13⁶$), 7.17–7.30 (10H, m, Ph), 7.40 (8H, 2 \times d, J=8.4 Hz, H-3['],5'), 7.94 (8H, 2 \times d, J= 8.3 Hz, H-2',6'); δ_C (75 MHz, CDCl₃) 14.0 (NCHCH₃), 20.3 and 20.5 ($-CH_3$), 21.6 (C-5,9), 21.7 (C-7^{*i*}), 30.8 (C-6,8), 31.9 and 32.1 (C-14,16,17,19), 46.3 (C-2,12), 47.6 $(-OCH_3)$, 48.7 (C-4,10), 57.1 (C- α), 103.0 (C-7), 108.7 $(C-1^3, 13^3)$, 115.1 $(C-15^5, 18^5)$, 118.6 and 119.1 $(C-1^1, 1^5)$ 13¹,13⁵), 123.6 (C-1⁶,13⁶), 127.1 (Ph),127.5 (C-15²,18²), 128.0 (Ph), 128.4, 128.5 and 128.6 $(C-2', 6' + Ph)$, 130.1 $(C_2^3, 5^1)$, 133.3 (C_1^1) , 139.9 and 140.1 $(C_1^1, 15^3, 18^1,$ 18³), 140.2 (Ph), 145.0 and 145.1 (C-15⁴, 15⁶, 18⁴, 18⁶), 145.6 $(C-4)$, 153.1 and 154.0 $(C-1^2, 1^4, 13^2, 13^4)$; MS: m/z (rel. int.) = 1584.1 $[M^+ + H]$ (4), 1520.9 (3), 1413.8 (3), 1185.5 (5), 1031.5 [-Tos] (3), 875.4 [–Tos] (3), 719.4 [–Tos] (3), 335.4 (70), 175.1 (67), 105.1 [PhCHCH3] (100). Found C, 65.98; H, 5.96; N, 1.76; S, 7.76%; $C_{87}H_{94}N_2O_{18}S_4$ requires C, 65.97; H, 5.98; N, 1.77; S, 8.10%.

6.2.14. (R,R) -1²,1⁴,11²,11⁴-Tetramethanesulfonyloxy-12, 14,15,17-tetramethyl-3,9-bis(1-phenylethyl)-13 4 ,13 6 ,16 4 , 16⁶ -tetra(p-toluenesulfonyloxy)-3,9-diaza-1,11(1,3,5), 13,16 (1,3)tetrabenzenabicyclo[9.3.3]heptadecaphane (15). Bridged resorcinarene 3a (450 mg, 0.3 mmol) and diisopropylethylamine (1.05 mL, 6 mmol) were dissolved in dry dichloromethane (15 mL). The solution was cooled to 0^oC in an ice-water bath and methanesulfonyl chloride (0.46 mL, 6 mmol) was added dropwise. The solution was then allowed to warm to room temperature. After the reaction was complete (tlc), it was poured into sat. NaHCO₃ (25 mL) and extracted with dichloromethane $(3 \times 25 \text{ mL})$ in the usual manner. After drying the organic layers with magnesium sulphate, and evaporation of the dichloromethane, the crude product was purified via column chromatography (40 g silica gel eluting with ethyl acetate–petroleum ether 1:1). The resulting material was then recrystallized (dichloromethane/ethyl acetate/petroleum ether) to give 15 (470 mg, 87%). Mp 194-195 °C (dichloromethane/ethyl acetate/petroleum ether); $[\alpha]_D$ -20.0 (c 2.9, CHCl₃); $\nu_{\text{max}}/\text{cm}^{-1}$ (CHCl₃): 3021m (aryl stretch), 2940s (C–H aliphatic), 1598m (aryl stretch), 1452s (CH₃), $1377s+1178s$ (-SO₂-O-); δ_H (400 MHz, CDCl₃) -0.95 (2H, m, H-6), -0.62 and -0.54 (4H, m, H-5,7), 0.89 and 0.93 (12H, $2 \times d$, $J=7.1$ Hz, $-CH_3$), 1.20 (6H, d, $J=6.8$ Hz, NCHC H_3), 1.35 and 1.52 (4H, m, H-4,8), 2.46 and 2.47 (12H, $2 \times s$, H-7[']), 3.24 (12H, s, $-SO_2CH_3$), 3.44 and 4.00 (4H, d, J_{AB} =14.7 Hz, H-2,10), 3.88 (2H, q, J= 6.8 Hz, NCHCH₃), 4.96 and 5.08 (4H, $2 \times q$, $J=7.1$ Hz, H-12,14,15,17), 6.84 (2H, s, H-13²,16²), 7.01–7.12 (10H, m, Ph), 7.36 and 7.39 (8H, $2 \times d$, $J = 8.5$ Hz, H-3', 5'), 7.44 (2H, s, $H-1⁶,11⁶$), 7.70 (2H, s, $H-13⁵,16⁵$), 7.91 and 7.97 (8H, d, $J=8.5$ Hz, H-2',6'); δ_C (75 MHz, CDCl₃) 13.7 (NCHCH₃), 21.2 (C-6), 21.4 (C-7'), 21.7 and 22.2 (-CH₃), 25.1 (C-5,7),

30.0 (C-12,14,15,17), 38.0 and 38.6 (-SO₂Me), 48.4 $(C_24, 8)$, 51.8 $(C_22, 10)$, 62.9 (NCHCH₃), 116.5 (C-13⁵, $16⁵$), 123.7 (C-1⁶,11⁶), 126.5 (C-13²,16²), 127.0 (Ph), 127.7 $(Ph), 129.2 (C-2', 6'), 130.0 (C-3', 5'), 132.6$ and $132.8 (C-1'),$ 134.8 (C-1³,11³), 136.6 and 137.2 (C-1¹,1⁵,11¹,11⁵,13¹, 13³,16¹,16³), 143.0 (Ph), 143.4, 144.3, 144.8 and 144.7 (C- 1^2 , 1^4 , 11^2 , 11^4 , 13^4 , 13^6 , 16^4 , 16^6), 145.9 (C-4'); m/z (rel. int.)=1807.9 $[M+H]$ ⁺(57), 1728.9 (21), 1653.0 (12), 510.4 (12), 105.1 [PhCHCH3] (100). Found C, 57.18; H, 5.31; N, 1.72; S, 14.05%; $C_{87}H_{94}N_2O_{24}S_8 \cdot H_2O$ requires C, 57.22; H, 5.30; N, 1.53; S, 14.05%.

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Enantioselective addition of diethylzinc to benzaldehyde catalysed by chiral, bridged resorcinarenes: a stereoselectivity model based on chirality transfer

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Abstract—The enantioselective addition of diethylzinc to benzaldehyde catalysed by a range of chiral bridged resorcinarenes has been studied, and the results used as a means of probing cooperative effects in the resorcinarene bowl. A structure–activity relationship has emerged in which bridged resorcinarenes with little available room in the bowl (e.g., **3b. 3c**) favour R-enantioselectivity in the product, while those promoting cooperative effects in the bowl via coordination sites in the bridge $(3e)$ or strong donor protecting groups $(3j)$ favour S-enantioselectivity. A mechanistic hypothesis based on Noyori's model to account for these trends has been put forward in which stereoselectivity is dependent on two factors as the ratio of axially diastereomeric anti-zincoxazines as well as the *exo* or *endo* bias of active zinc. The model explains why ee's are never greater than around 50% and indicates asymmetric induction to be due to the axial chirality in the bowl as a result of chirality transfer from the bridge, rather than due to induction via the central chirality in the line of the bridge. As a result, the model stimulates some new fascinating possibilities in enzyme mimicry.

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1. Introduction

In the previous paper, 1 we described the synthesis and characterization of a range of chiral bridged resorcinarenes using Mannich technology for bridge incorporation. Emphasis was placed on varying the protecting groups on the upper rim as well as the length and nature of the chiral diamine line used to construct the bridge. Although a range of pendant lower-rim groups were screened in the simple model, only $R = CH_3$ was used for the full set of derivatives. Our purpose for synthesizing these derivatives was to investigate the potential use of such structures as templates for asymmetric catalysis. The literature contains very few examples of calixarenes and resorcinarenes in asymmetric catalysis. For instance, Matt has shown that a lower-rim, inherently chiral calixarene scaffold can be used in allylic alkylation (palladium) and hydrogenation (rhodium), although low ees were obtained. 2 Others have shown that cooperative effects may potentially be provided by supramolecular interactions involving the concave bowl.^{[3](#page-162-0)} Of particular interest for us was the possibility of using cooperative effects, which could be investigated using an asymmetric reaction as a probe. Given the presence of tertiary amino and phenolic hydroxyl functionality in a 1,3-relationship near to the chiral auxiliaries in the bridge, it was decided to use the enantioselective addition of diethylzinc to benzaldehyde as this probe reaction, given that this reaction is mechanistically well-documented and known to work with γ -amino alcohols.^{[4,5](#page-162-0)}Many catalysts based on chiral β - and γ -amino alcohols have been reported for the enantioselective addition reaction of diethylzinc to benzaldehyde (aldehydes), with many of them now able to achieve ee's in excess of 95% .^{[6](#page-162-0)} A comprehensive mechanistic picture has now emerged in which the amino alcohol functionality is postulated to form a zincoxazine intermediate that acts as a template for catalysis.^{[7](#page-162-0)}

To this end we have recently reported evidence for the first example of a cavity-controlled addition of diethylzinc to benzaldehyde involving ketal functionality in the bridge. 8 In 8 In this paper, we present results on a comprehensive study of the diethylzinc reaction using our bridged resorcinarenes as catalyst templates, and use the trends in stereoselectivity to develop a mechanistic model that offers insights into the origin of the enantioselectivity, and hence ideas on future design.

Keywords: Resorcinarene; Catalytic template; Diethylzinc; Enzyme mimicry; Cavity control.

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Scheme 1. Synthesis of bridged resorcinarenes. Reagents and conditions: (i) $(CH₂O)_n$, CH₃CN, 90 °C.

2. Results

Our first studies were conducted on bridged resorcinarene tetratosylates $3a-e$ ^{[8](#page-162-0)} These could be synthesized by the Mannich reaction between resorcinarene tetratosylate 1, diamine 2 and excess paraformaldehyde in moderate to good yields (Scheme 1, Table 1), 8.9 in which the resorcinarene tetratosylates 1 could themselves be syn-thesized by known methodology.^{[9–11](#page-162-0)} The derivatives chosen aimed to firstly probe communication between lower and upper-rims by varying the pendant R group, while keeping the bridge constant, and secondly to probe the influence of having a coordinating (to zinc) group (X) in the middle of the bridge while keeping the pendant R group constant (as methyl).

Table 1. Yields for bridged resorcinarenes synthesized

	R	Х	Yield $(\%)$
3a	CH ₃	Н	60
3 _b	C_5H_{11}	Н	80
3c	$C_{11}H_{23}$	Н	67
3d	CH ₃	$OCH2CCH3)2CH2O$	47
3e	CH ₃	OMe	57

Addition reactions were carried out in all cases using benzaldehyde as limiting reagent with diethylzinc (2 equiv) and a resorcinarene loading of 5 mol% in toluene at room temperature for 24 h. Yields of 1-phenyl-1-propanol were ascertained from purification using column chromatography while ee's were determined by chiral GC using a modified b-cyclodextrin glass capillary column (Heptakis-2,3-di-Oacetyl-6- O -TMBMS- β -CD). The results can be found in Table 2.

Two important trends emerge from these results. The first was that the enantioselectivity increased with lengthening of the resorcinarene pendant R groups, which drew attention to the possible involvement of the cavity in influencing the stereochemical outcome of the reaction. This was because ¹H NMR spectroscopic studies on resorcinarenes 3a-c, as described in a previous paper, $¹$ $¹$ $¹$ had revealed an increasing</sup> downfield shift of the bridge methylenes $(X=H \text{ in } 3)$ going from 3a to 3c, indicating that the bridge spent more time in

the cavity for $3c$.^{[12](#page-162-0)} This trend was rationalized by invoking increased lower-rim repulsions for 3c resulting in the upper rim phenyl groups being pushed closer together. Importantly for these results, the increase in ee on going from 3a to 3c would thus be associated with minimizing room in the bowl, which has implications for the mechanistic model proposed later. The second trend identified, intriguingly, was that the dimethoxy ketal on the bridge of 3e effected a complete reversal of the enantioselectivity of the reaction. We have recently proposed that this result suggests intracavity catalysis in view of the position of the ketal in the middle of the line and the possibility therefore of direct involvement of the ketal group as a coordination site to zinc.[8](#page-162-0)

Table 2. Results for the asymmetric addition of diethylzinc to benzaldehyde

		cat (5 mol%) Et ₂ Zn (2eq) PhCH ₃ /rt 24h		ЭH
	R	X	Yield $(\%)$	ee $(\%)$ (enantiomer)
3a 3b 3c 3d 3e	CH ₃ C_5H_{11} $C_{11}H_{23}$ CH ₃ CH ₃	Н H Н $OCH2CCH3)2CH2O$ OMe	91 94 99 85 85	12(R) 41 (R) 52 (R) 34(R) 51 (S)

Satisfied that the enantioselective addition of diethylzinc to benzaldehyde could indeed be used as a mechanistic probe reaction, we elected to broaden the study to other structural changes, notably to address the influence of the upper-rim protecting groups as well as the length of the bridge. These changes were addressed in that order.

Alternatives to the tosylate 'protecting' group were selected for ease of synthesis and potential for providing different steric and electronic environments near the cavity. Their synthesis and characterization has been described in a previous paper.^{[1](#page-161-0)} Thus, methanesulfonate (OMs, $3f$) and triisopropylbenzenesulfonate (OTrips, 3g) were chosen to sterically complement the p-toluenesulfonate (OTs, 3a)

groups already used. In addition, ester derivatives such as acetate (OAc, 3h) and tolyl (OTol, 3i) were synthesized, along with a benzyl carbonate (OCBz, 3j) in order to investigate electronic effects around the group directly attached to the phenolic oxygen. Attempts at synthesizing an alkyl methyl analogue failed. Slightly longer diamine lines for bridging were synthesized and characterized, including an example with a dimethoxy ketal in the centre of the bridge (3l) for comparison with the shorter-bridged resorcinarene 3e. In all these cases the pendant R group was kept as methyl.

These new compounds were then evaluated out as before for their ability to catalyze the asymmetric addition of diethylzinc to benzaldehyde. The results are shown in Table 3, along with a comparison with compound 4, an o-hydroxybenzylamine containing a chiral a-methylbenzyl auxiliary at nitrogen, which was prepared by Palmieri and investigated in the same reaction.^{[13](#page-162-0)} Compound 4 modelled a structural fragment of bridged resorcinarene 3.

Table 3. Results for the asymmetric addition of diethylzinc to benzaldehyde

^a Benzyl alcohol (46%) obtained as byproduct.

^b Benzyl alcohol (91%) obtained as byproduct.

Table 3 reveals that the ee's showed no quantitative improvement compared to those obtained previously. An important observation though was that the results with 4 indicated that the resorcinarene skeleton does indeed influence the enantioselectivity. Palmieri found that the best results ($>90\%$ ee) were obtained when a chiral group was attached to position 1 (marked on table diagram) indicating that the resorcinarene structure plays a pivotal role in the asymmetric induction.

The trend in ee's provided an opportunity to draw some interesting conclusions. In this regard, the derivatives should be viewed as four sets as: 3f/3g; 3h–3j, 3k and 3l. Firstly, the 3f/3g comparison: increasing the steric bulk of the sulfonate protecting group from tosylate (3a, [Table 2](#page-155-0)) to the triisopropylbenzenesulfonate (Trips, 3g), did not

significantly alter the reaction stereo-outcome. However, reducing its size to the methanesulfonate (Ms, 3f), resulted in a reversal in the sense of enantioselectivity in the product towards favouring the S-enantiomer. In the 3h–3j set, changing the protecting group from a sulfonate to an ester (Ac, Tol) or carbonate (Cbz), resulted in the product always favouring the S-configuration rather than $R₋$, with the carbonate returning the best and significantly higher ee for the unfunctionalised $(X=H)$ bridged resorcinarenes. Thirdly, increasing the length of the unfunctionalised bridge by two methylene groups as 3k while keeping tosylate as protecting group also resulted in a reversal of enantioselectivity from R to S . Lengthening the functionalised bridge of 3e by two methylene groups to 3l resulted in retention of the S-enantioselectivity but at a reduced level (51 vs 21%).

In our previous work focusing on compound $3e$, we suggested that a transition-state occurring in the resorcinarene cavity equated with reversal of enantioselectivity. Thus, in accordance with this hypothesis, the results above would be rationalised as follows. Firstly, changing from the larger tosylate to the smaller mesylate group decreases steric bulk near the cavity and thus allows for a greater chance of an intracavity transition-state. Secondly, changing from sulfonate to ester protecting groups (i.e., from tetratosylate 3a to tetratolate 3i), results in oxygen donor atoms near the cavity with increased Lewis basicity, thus directing the diethylzinc into the cavity (see Fig. 1). This is particularly poignant for the tetrabenzylcarbonate 3j, which returned an ee of 49% (S), as a result of having two mesomeric oxygen donors to the carbonyl group.

Figure 1. Relative donor strengths of protecting group oxygens.

Similarly for the longer-bridged resorcinarene 3k, the selectivity shift from R to S was also indicative of more available room in the cavity in keeping with the NMR trends.^{[1](#page-161-0)} The upfield shift for the longer bridge $3k$ was less $(-0.22$ ppm) compared to that of the shorter bridged 3a $(-0.37$ ppm) indicating that the bridge of 3k lies on average more out of the cavity compared to 3a, probably to maximize entropy. Furthermore, the longer-bridged 3l containing the ketal in the centre of the bridge also returned an S-enantioselectivity indicating a predominantly cavitycontrolled process. However, the ee was lower than that of the shorter bridged compound $3e(21 \text{ vs } 51\%)$ presumably as a result of the ketal coordinating group being on average further out of the cavity. All of these results appeared to corroborate our hypothesis for 3e put forward to explain the trends in enantioselectivity as S-enantioselectivity indicating a preferred cavity-controlled process. A further important observation made from the results, in particular

for the longer-bridged resorcinarenes 3k and 3l, was the reduction in chemical yield compared to other bridged resorcinarenes. For 3k and 3l, the byproduct produced in these reactions was determined to be benzyl alcohol, produced via hydride reduction of the substrate benzaldehyde. Ridley and Coates, in their study of organozinc compounds, have proposed that diethylzinc can do this via a b-elimination of hydride from diethylzinc with concomitant release of ethylene.^{[14](#page-162-0)} This was supported by the observation that the reaction of diethylzinc with benzophenone gave the reduced product quantitatively, whereas no reaction (addition or reduction) was observed with dimethylzinc, which lacks β -hydrogens. The authors proposed a cyclic transition-state shown in Figure 2.

$$
\begin{array}{ccc}\n & H \\
 & \searrow & \searrow \\
 & \searrow & \searrow & \searrow \\
 & H \\
 & \searrow & \searrow & \searrow \\
 & H \\
 & H \\
 & H\n\end{array}
$$

Figure 2. Proposed six-membered transition-state for reduction of benzophenone.

In the case of 3k and 3l, presumably the longer and hence larger bridges make it energetically more difficult to accommodate the benzaldehyde substrate inside the cavity. The result is that the diethylzinc favours the less sterically demanding hydride delivery option compared to the ethyl transfer. This scenario is particularly significant for ketal 3l in which the ketal attempts to direct zinc coordination and hence reaction from within the cavity of the bowl, but the larger line promotes a favoured hydride delivery on steric grounds.

3. Mechanistic model

In order to consolidate our thoughts into a mechanistic model, attention was turned towards the current mechanistic view on this reaction. Of particular significance for our results was the observation that ee's were never higher than around 50%, which made us uncomfortable when referring to a 'cavity-controlled' process. Thus, we considered this feature to be highly significant. Although several reaction mechanisms involving various ligands have appeared in the literature, that of Noyori's utilizing amino alcohols as ligands^{[7](#page-162-0)} has received the widest recognition. This mechanism postulates that two molecules of dialkylzinc per catalyst are required for a reaction to occur, the first forming a zincchelate complex with expulsion of alkane to provide a zincoxazine template (template-zinc) for the complexation of the second equivalent that generates the active zinc-entity (active-zinc) responsible for alkyl group transfer. Palmieri,[8](#page-162-0) working on derivatives of 4 as catalysts for the same reaction, has also postulated an analogous 'zincoxazine' template formation with dimethylzinc, based on Noyori's work. The full mechanism is depicted in Figure 3.

For mechanistic studies regarding the ratio of diethylzinc to the pre-catalyst resorcinarene, a series of reactions were conducted a using a 1:1 ratio of catalyst: benzaldehyde, so as to be able to determine the number of equivalents of diethylzinc required for the reaction to go to completion. Two resorcinarenes were studied on a 0.5 mmol scale, 3b $(R=n-pentyl)$ and 3e $(R=CH_3)$ with the dimethoxy ketal in the bridge. The pendant R group and the group in the bridge were varied in this study in order to probe their importance. Moreover, carrying out the reactions on a 0.5 mmol scale

Figure 3. Palmieri's template and active zincoxazines.

minimized errors of judgment that might be incurred in the syringing of the reactants. Reactions were left the customary 24 h after the addition of each equivalent of diethylzinc to allow the possibility of reaction to proceed before monitoring by TLC. The results are shown in Table 4.

Table 4. Equivalents diethylzinc required to ensure complete reaction

$Et2Zn$ (equiv)	Resorcinarene 3b	Resorcinarene 3e
	No reaction	No reaction
	Partial ^a	No reaction
	Complete reaction	No reaction
		Complete reaction

^a Judged by TLC to be \sim 10%.

With a ratio of $2:1:1$ Et₂Zn/resorcinarene/benzaldehyde, neither resorcinarene produced any reaction. At 3:1:1, 3b showed partial reaction (\sim 10% by TLC), while 3e showed no conversion of starting material. At 4:1:1, 3b showed complete conversion to the product 1-phenyl-1-propanol, while 3e remained at 0% conversion, Addition of the fifth equivalent of diethylzinc to 3e finally resulted in complete conversion of benzaldehyde. These trends were considered to be highly significant indicators for constructing a mechanistic model, which we now delineate.

In keeping with the Noyori mechanistic model, these results support the notion of initial consumption of 2 equiv of diethylzinc to form a bis-zincoxazine template intermediate, for which theoretically there are three possible arrangements, as one syn and two axially diasterisomeric antizincoxazine templates (Fig. 4).

The third equivalent of diethylzinc could either coordinate to a zincoxazine oxygen to form 'active-zinc' (see [Fig. 3\)](#page-157-0), resulting in alkyl addition to benzaldehyde (once coordinated), or it could react with one of the unreacted, poorly nucleophilic hydroxyl groups, resulting in an inactive site in view of the inability to form a chelate. Figure 5 shows three possible pathways that might exist for the reaction of

Figure 4. The three possible arrangements of the bis-zincoxazine resorcinarene templates.

svr

Figure 5. Possible mechanism for coordination of third diethylzine molecule.

Figure 6. Possible mechanism or the coordination of third and fourth diethylzine molecules in 3e.

Figure 7. 3D-representation of endo versus exo facial delivery for zincoxazine in the anthoclockwise configuration (showing only one of the zincoxazines and including coordinated 'active-zinc').

the third equivalent of diethylzinc. Mechanistically, these suggest an initial coordination of the diethylzinc to the oxygen of one of the zincoxazines, followed by an intramolecular rearrangement of the 'active-zinc' onto a free hydroxyl group either on the opposite side of the bowl (pathway A) or on the same side (pathway B). This complexation offers a plausible mechanism for the interconversion of the *syn/anti*-bis-zincoxazine arrangements as shown by pathway C. Evidently a competition exists between active-zinc formation and hydroxyl exchange, with the latter being favoured, since the third equivalent of diethylzinc resulted in only partial conversion of benzaldehyde to 1-phenylpropanol for 3b and none for 3e. It was only on addition of the fourth equivalent of diethylzinc that the 3b reaction went to completion, now indicating a greater preference for activation of 'templateddiethylzinc' to form 'active-zinc' compared to reaction with the final free hydroxyl group.

For 3e, however, conversion was still blocked, either as a result of exchange with the remaining phenolic hydroxyl or because of competing coordination of diethylzinc by the dimethoxy ketal. We consider the latter to be more plausible, particularly in view of the results with 3b. Finally, addition of the fifth equivalent to the 3e reaction successfully brought about full conversion indicating that active-zinc is only set up after the fourth equivalent of

diethylzinc with this template. Figure 6 shows how these processes might possibly occur.

The next step in the Noyori model requires coordination of the benzaldehyde oxygen to the zinc of the templated-zinc. In our case it seems reasonable to postulate that the benzaldehyde approaches the zincoxazine in a preferred orientation governed by steric interactions, regardless of inter- versus intracavity reaction preference. Consideration of molecular models suggests this to involve an orientation in which the phenyl group points away from both the bridge and the lower rim. Once both 'active-zinc' has been set up and benzaldehyde coordination to 'templated-zinc' has taken place, ethyl group transfer may proceed from the *exo*- or *endo*-face of the zincoxazine,[†] depending on the positioning of 'active-zinc' endo-face attachment of 'active-zinc' is shown in Figure 7.

Of the three diastereoisomeric bis-zincoxazines depicted in [Figure 4,](#page-158-0) literature precedent suggests that the antiarrangement predominates. Böhmer has demonstrated that Mannich condensation of resorcinarene tetratosylates (1) with achiral primary diamines and formaldehyde gave predominantly bridged anti bis-benzoxazines with

 \dagger The endo-face is synonymous with the cavity, while the exo-face effectively places the active diethylzinc on the outside of the perimeter.

Scheme 2. Reagents and conditions: (i) (R) - α -methylbenzylamine, formaldehyde, AcOH (cat.), EtOH, rt.

the syn-isomer only being detected by ¹H NMR (<5%) when the length of the polymethylene spacer is five or more carbons \log ^{[15](#page-162-0)} Further support for discounting the syndiastereomer comes from our own work in which condensation of resorcinarene tetratosylate 1 with chiral (R) - α -methylbenzylamine and formaldehyde gave a 3:1 mixture of the anti bis-benzoxazine 5 with no trace of the syn-isomer, Scheme 2. The ${}^{1}H$ NMR spectrum (Fig. 8) showed three sets of aromatic methine singlets in the 6.2–7.0 ppm range, which is characteristic of the antibisbenzoxazine arrangement of a mixture of diastereomers (the syn-arrangement would have resulted in the appearance of five signals in this region).

Thus, in this transition-state model the enantioselectivity of the reaction is determined by the interplay between the exoversus endo-delivery bias of 'active-zinc', together with the diastereomeric excess of the two anti bis-zincoxazines, which together account for the large range of ee's from R to S. It is likely that the latter is primarily due to the exo/endo factor rather than a shift in the anti bis-zincoxazine equilibrium, as the exo/endo bias would presumably be more responsive towards structural changes affecting the space in the cavity, and hence the positioning of active zinc inside or outside. Consequently, this model indicates that enantioselectivity is not as a result of the $sp³$ $sp³$ $sp³$ asymmetric chirality in the bridge but rather due to axial chirality in the bowl^{[16](#page-162-0)}—as a result of chirality transfer from the bridge working in conjunction with cavity effects. Such a model neatly explains why ee's are never greater than 50%. For instance, the ketal-bridged template 3e result, with an ee of 51% in favour of the S-enantiomer, may be rationalized as proceeding via two axially diastereomeric zincoxazines, one giving rise to 75% S and the other 25% R, but both via *endo*delivery of ethyl to a benzaldehyde positioned in both cases with its phenyl group pointing away from the resorcinarene.

Similarly, our hypothesis that S-enantioselectivity is synonymous with endo delivery of the ethyl group, allows prediction that the dominant bis-zincoxazine diastereomer of the 3:1 mixture is the anticlockwise one. This correlates with Heaney's work on resorcinarene tetrabenzoxazines in which the *R*-auxiliary promoted a bias to the anticlockwise isomer.^{[17](#page-162-0)}

The present transition-state model may be thus summarised as follows:

- Reaction proceeds via two axially-chiral, diastereomeric anti bis-zincoxazines, whose ratio is estimated to be 75:25 in favour of the 'anti-clockwise'— isomer when the bridge is (R, R) . This would explain why ee's were never greater than about 50%.
- The stereo-positioning of active zinc as either *exo* or endo with respect to the cavity (bowl) is a crucial feature in determining R or S enantioselectivity. Delivery of ethyl from exo-'active-zinc' is considered to produce an excess of the R-enantiomer of 1-phenyl-1-propanol. Thus, compounds 3b and 3c with long pendant chains and minimised room in the bowl set up a predominance of exo-active zinc, which results in exo-delivery of ethyl favouring the R-products.

Figure 8. ¹H NMR spectrum of 5 showing the major and minor isomers.

• Conversely, *endo*-attachment of active zinc resulting in endo delivery of ethyl favours the S-configuration in the product. This is the case for compounds 3e and 3j involving cooperative cavity effects—the former from donor methoxy groups in the bridge, and the latter from the electron-donating carbonate groups—which promote the kinetic formation of endo-'active-zinc'.

4. Conclusion

The enantioselectivities achieved by these catalysts for the reaction in question by no means compare with the high values ($>95\%$ ee) achievable for many other systems.^{[4](#page-162-0)} However, this study has allowed generation of a working model alluding to some interesting cavity effects operating in conjunction with axial chirality in the resorcinarene bowl, as a result of chirality transfer from the asymmetry in the bridge, as being responsible for the enantioselectivity. The study^{[8,9](#page-162-0)} represents the first reported example of using the bowl of an asymmetrically functionalised resorcinarene for promoting asymmetric catalysis, 8 a desirable research objective suggested recently by Iwanek.[18](#page-162-0) In this regard, these catalysts open up a new dimension in resorcinarene asymmetric catalysis and synthetic enzyme mimicry.

5. Experimental

5.1. General remarks

All diethylzinc reactions were carried out under argon using dry toluene. Nuclear magnetic resonance spectra were recorded on a Varian Unity 400 (100 MHz for 13 C) or Varian Mercury 300 MHz (75 MHz for 13 C) and were carried out in chloroform-d. Optical rotations were obtained using a Perkin Elmer 141 polarimeter at 20° C. Melting points were obtained using a Reichert Jung Thermovar hotstage microscope and are uncorrected. Elemental analyses were performed using a Fisons EA 1108 CHN elemental analyser. Infrared spectra were recorded on a Perkin-Elmer Paragon 1000 FT-IR spectrometer in either dichloromethane or chloroform. Enantiomeric excesses were determined by chiral GC on a Hewlett Packard HP 5890 Gas Chromatagraph, using a modified β -cyclodextrin glass capillary column (Heptakis-2,3-di-O-acetyl-6-O-TMBMS- β -CD) and helium as a carrier gas. Bridged resorcinarenes 3 were synthesised according to reported procedures.^{1,8}

5.2. General procedure for the synthesis of 1-phenylpropanol

To a pre-dried Schlenk tube was added catalyst (0.025 mmol) and 1 M diethylzinc in toluene (1 mmol, 1 mL) at -20 °C. The solution was allowed to stir for 15 min followed by the addition of 1 M benzaldehyde in toluene (0.5 mmol, 0.5 mL). The reaction vessel was then sealed and allowed to warm slowly to room temperature. After 16 h the reaction was checked (TLC) for complete consumption of benzaldehyde. On completion, the reaction was cooled to 0° C and quenched by the slow addition of 1 M HCl (1 mL), and then extracted from water (20 mL) with dichloromethane $(3 \times 20 \text{ mL})$. The organic extracts

were dried over magnesium sulphate and reduced in the usual manner. Flash chromatography $(5 g SiO₂, eluting)$ with ethyl acetate/petroleum ether 3:17) afforded 1-phenylpropanol as a clear oil. The enantiomeric excess was determined by chiral GC.

5.2.1. $(R,R)-1^5,5^5$ -Dihydroxy-2,4,6,8-tetramethyl-1³, 5^3 -bis(1-phenylethyl)-3⁴,3⁶,7⁴,7⁶-tetra(p-toluene-sulfonyloxy)-3,7 (1,3)benzena-1,5(6,8)-(3,4-dihydro-2H-benzo[e]- [1,3]oxa-zina)cyclooctaphane (5). Tetramethylresorcinarene tetratosylate (290 mg, 0.25 mmol) and (R)- α -methylbenzylamine (0.2 mL, 1.5 mmol) were added to ethanol (30 mL) and 32% aqueous formaldehyde (8 mL). Glacial acetic acid (0.5 mL) was added and the reaction allowed to stir at room temperature overnight. The product that precipitated was filtered and washed with ethanol to give the title compound (280 mg, 80%) as an off-white powder. Mp 152–155 °C (from ethanol–dichloromethane); $[\alpha]_{\text{D}}$ –6.7 (c 1.67 in CHCl₃); $\nu_{\text{max}}/\text{cm}^{-1}$ (CHCl₃) 3514s $(O-H, H-bonded), 2930s + 2853s (C-H), 1598m + 1483m$ (aryl stretch), $1371s+1186s$ (–SO₂–O–); $\delta_{\rm H}$ (300 MHz, CDCl₃) 1.26–1.34 (6H, m, C-4/8 –CH₃), 1.40 (6H, d, J= 6.6 Hz, NCHC H_3), 1.43–1.50 (6H, m, C-2/6 –C H_3), 2.45, 2.46, 2.48 and 2.50 (12H, $4 \times s$, H-7[']), 3.36 and 3.80 (4H, $2 \times d$, $J_{AB} = 17.6$ Hz, $H - 1^4$, 5^4 -major), 3.77 and 4.02 (4H, $2 \times d$, $J_{AB} = 17.6$ Hz, H-1⁴,5⁴ -minor) 3.80 (2H, q, $J=$ 6.6 Hz, NCHCH₃), 4.09 and 4.67 (4H, $2 \times d$, $J_{AB} = 8.9$ Hz, $H-1²,5²$ -minor), 4.18–4.76 (4H, 2×d, $J_{AB} = 10.3$ Hz, H-1²,5² -major), 4.33 (2H, q, $J=6.8$ Hz, H-4,8 -major), 4.52 (2H, q, $J=6.8$ Hz, H-2,6 -major), 4.25 and 4.45 (4H, m, H-2,4,6,8 -minor), 5.85 (2H, s, –OH -major), 6.00 (2H, s, –OH -minor), 6.27 (2H, s, H-3² -major), 6.32 (2H, s, H-3² -minor), 6.54 (2H, s, $H-3^5$ -minor), 6.57 (2H, s, $H-3^5$ -major), 6.96 (2H, s, H-1⁷ -major), 6.98 (2H, s, H-1⁷ -minor), 7.18–7.47 (18H, m, Ph $\text{ + H-3/5'},$ 7.76–7.94 (8H, m, H-2 $\frac{1}{6}$; δ_C (75 MHz, CDCl₃) 19.1, 20.5 (CH₃ major), 20.6, 20.9 (CH3 minor), 31.4 (C-4,8 minor), 31.7 (C-4,8 major), 32.2 (C-2,6 minor), 32.4 (C-2,6 major), 43.8 (C-1⁴ minor), 45.4 (C-1⁴ major), 58.4 (NCHCH₃), 78.6 (C-1² major), 79.8 (C-1² minor), 108.9 (C-1¹⁰ major), 109.0 $(C^{-1}$ ¹⁰ minor), 114.6 (C-3⁵ minor), 114.8 (C-3⁵ major), 116.7 (C-1⁶ minor), 116.9 (C-1⁶ major), 121.3 (C-1⁸ major), 121.6 (C-1⁸ minor), 122.2 (C-1⁷ major), 122.3 (C-1⁷ major), 126.6 (C-3² major), 126.9, 127.2, 127.6, 128.0, 128.1, 128.3, 128.4, 128.6, 128.7, 129.8, 130.3, 130.4 (Ph + C-2[']/ $3¹ / 5¹ / 6¹$, 131.5 + 133.8 (C-1¹ minor), 131.6 + 134.1 (C-1¹) major), 138.2 (C-3¹ minor), 138.3 (C-3¹ major), 141.7 (C-3³) minor), 142.1 (C-3³ major), 143.9 (C-3⁶ minor), 144.2 (C-3⁶) major), 144.3, 145.2 + 145.3 (C-4^{/}+ Ph), 145.6 (C-3⁴ major), 146.3 (C-4'), 150.3 (C-1⁵ minor), 150.7 (C-1⁵) major), 151.8 (C-1⁹ minor), 152.3 (C-1⁹ major); MALDI TOF: m/z (rel int.) 1450.51 [M⁺ - H] (5); Found: C, 65.95; H, 5.41; N, 1.94; S, 8.34%; $C_{80}H_{78}N_2O_{16}S_4$ requires C, 66.19; H, 5.42; N, 1.93; S, 8.83%.

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Efficient synthesis of 18–40 membered macrocyclic polyoxadiamides and polyoxatetraamides via ring closing metathesis

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Abstract—Ring closing metathesis of the appropriate $1,\omega$ -dienes led to efficient synthetic approaches towards the corresponding macrocyclic polyoxadiamides and tetraamides with 18–40 membered ring sizes in good to excellent yields using Grubbs' catalyst. Q 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Recently, ring closing metathesis $(RCM)^1$ $(RCM)^1$ has found increasing application in the synthesis of macrocyclic compounds.²⁻¹² The application of RCM as the key macrocyclization step in the synthesis of crown compounds has opened efficient routes towards macrocyclic crown compounds of important applications in supramolecular chemistry.^{6,13-16}

The present work describes new routes for the synthesis of macrocyclic polyethers containing diamide and tetraamide groups inside the macrocyclic ring via RCM using Grubbs'catalyst I as the key macrocyclization step. The results obtained provide an efficient synthetic procedure to the macrocyclic polyoxadiamides 3b–e, 4, 5, 6a,b, 7a,b, 9a,b with 18–34 membered ring sizes and the macrocyclic polyoxatetraamides 1a–e, 2b,d, 8a,b with 28–40 membered ring sizes. The new macrocyclic crown-amides 1–9 synthesized in the present investigation are of potential interesting diverse applications, which have been reviewed in our recent publications (Fig. 1).^{[16](#page-176-0)}

2. Results and discussion

[Scheme 1](#page-165-0) illustrates our synthetic routes towards the precursors $1,\omega$ -dienes **15a–e** needed for the RCM synthesis of $1a-e$. Thus, acylation of o -aminophenol 10 with

 o -allyloxybenzoyl chloride 11 afforded o - $(o$ -allyloxybenzamido)phenol 12. The latter was converted into its potassium salt 13, which was then reacted with the appropriate bis-chloroacetanilide derivatives 14a–e to afford the corresponding $1,\omega$ -dienes **15a–e** in 63–93% yields. RCM of 15a–e proceeded smoothly to give the corresponding macrocycles 1a–e in 52–97% yields with Grubbs' catalyst I (2.5–5 mol%) in refluxing CH_2Cl_2 for 24 h. The results are presented in [Table 1](#page-165-0).

On the other hand, $1,\omega$ -dienes 17b,d were readily obtained as outlined in [Scheme 2.](#page-165-0) Thus, reacting the potassium salt of o-allyloxyacetamidophenol 8 with each of 14b,d afforded the corresponding desired $1,\omega$ -dienes 17b,d. RCM of the latter dienes proceeded smoothly to give the corresponding macrocycles 2b,d in 70 and 60% yields, respectively, upon heating with Grubbs' catalyst I (5 mol%) in refluxing CH_2Cl_2 for 24 h. The results are presented in [Table 1.](#page-165-0)

[Scheme 3](#page-166-0) outlines the synthetic routes followed for the synthesis of the $1,\omega$ -dienes 21–23, required for the RCM synthesis of macrocycles 3b–e, 4 and 5. Thus, alkylation of methyl salicylate 18 with 2-allyloxyethyl p-toluenesulfonate afforded methyl 2-(2-allyloxyethoxy)benzoate, which was hydrolysed to the corresponding acid. The latter was then converted to the corresponding acid chloride 19 by the action of $S OCl₂$. Reaction of 19 with each of the amines 20b–e, ethylenediamine and o-phenylenediamine afforded the corresponding $1,\omega$ -dienes 21b–e, 22 and 23, respectively. Compounds 22 and 23 were also, prepared by reacting the dipotassium salts of each of 1,2-bis(2-hydroxybenzamido)ethane and 1,2-bis(2-hydroxybenzamido) benzene with 2-allyloxyethyl p-toluenesulfonate. RCM of

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Figure 1.

compounds 21–23 afforded corresponding macrocyclic compounds 3–5 in 60–99% yields upon heating with Grubbs' catalyst I (2.5–5 mol%) in refluxing CH_2Cl_2 for 4–24 h. The results are presented in [Table 2](#page-166-0).

On the other hand, $1,\omega$ -dienes 30–33 were readily obtained as outlined in [Scheme 4](#page-167-0). Thus, acylation of each of the o -allyloxyaniline 24a and o -(2-allyloxyethoxy)aniline 24b with chloroacetyl chloride afforded the corresponding chloroacetanilide derivatives 25a,b, respectively. Reacting the latter with the potassium salt of the appropriate phenolic compounds 26–29 afforded the corresponding desired $1, \omega$ -dienes 30–33. RCM of the latter dienes proceeded smoothly to give the corresponding macrocycles 6–9 in 73–100% yields upon heating with Grubbs' catalyst I $(2.5-7.5 \text{ mol\%)}$ in refluxing CH₂Cl₂ for 2–24 h ([Table 3](#page-167-0)).

In all RCM reactions, the progress of the reaction was monitored by TLC and ¹H NMR analysis where no further

increase in products was noticed (after the time indicated in [Tables 1–3](#page-165-0)) of reflux in CH_2Cl_2 . The RCM products 1–9 were shown from their ¹H and ¹³C NMR to consist of the E and Z isomers in different ratios as shown in [Tables 1–3](#page-165-0).

The E:Z isomers of the olefinic crown-diamides were readily assigned and their ratios were determined from the ¹H and 13 C NMR spectra. The major products in all RCM reactions were shown to be the E isomers with the characteristic 13 C signal of the OCH₂ (of the $OCH₂CH=CHCH₂O$, which appears further downfield than that for the corresponding \overline{Z} isomer. In ¹H NMR the OCH_2CH and OCH_2CH signals of E isomers appear as broad singlets. On the other hand, those similar proton signals for the Z isomer split into a doublet and triplet with $J=3-5$ Hz.

The present work describes an efficient synthetic access towards macrocyclic crown-diamide and crown-tetraamide

Scheme 1.

Table 1. Catalyst %, yields and E/Z ratios of macrocycles 1a-e, 2b,d

Entry	Substrate	(mol%) Catalyst/substrate, reaction time (h)	Yield (%)	Product $E:Z$ ratio
-1	15a	2.5, 24	97	$1a\,5:1$
$\mathbf{2}$	15 _b	5.0, 24	91	$1b$ 7:1
3	15c	2.5, 24	52	1 c 5:1
$\overline{\mathbf{4}}$	15d	2.5, 24	78	1d $5:1$
5	15e	2.5, 24	88	1e $13:1$
6	17 _b	5.0, 24	70	$2b$ 2.5:1
7	17d	5.0, 24	60	$2d$ 3.5:1

derivatives, with potential diverse applications in supramolecular chemistry and as starting compounds for further synthetic transformations utilizing the RCM techniques for the macrocyclization step. The examples of RCM presented here represent one of the best macrocyclization reaction techniques for the synthesis of crown compounds. It also, expands the utility of RCM methodology and its application to the synthesis of cyclic olefins of large ring sizes with different functional groups.

3. Experimental

3.1. General

All melting points are uncorrected. IR spectra were recorded in KBr disks using Perkin Elmer System 2000 FT-IR spectrophotometer. ^IH and ¹³C NMR spectra were recorded on a Bruker DPX 400, 400 MHz super-conducting NMR spectrometer. Mass spectra were measured on VG Autospec-Q (high resolution, high performance, tri-sector GC/ MS/MS) and with LCMS using Agilent 1100 series LC/ MSD with an API-ES/APCI ionization mode. Microanalyses were performed on LECO CH NS-932 Elemental Analyzer. The starting o -allyloxybenzoyl chloride 11,^{[17](#page-176-0)} bis-amines $14a-e$, ^{[16e](#page-176-0)} bis-amides $20b-e$, ^{[18](#page-176-0)} 1,2-bis(2-hydroxybenzamido)ethane^{[19](#page-176-0)} and 1,2-bis(2-hydroxybenz-amido)benzene^{[19](#page-176-0)} were prepared as reported.

3.1.1. 2-(o-Allyloxybenzamido)phenol 12. To an ice cold $(0-5 \degree C)$ stirred solution of 10 (5.3 g, 49 mmol) and

Scheme 3.

Table 2. Catalyst %, yields and E/Z ratios of macrocycles 3b–e, 4, 5

Substrate	(mol%) Catalyst/substrate, reaction time (h)	Yield $(\%)$	Product $E:Z$ ratio
21 _b	2.5, 24	99	$3b \; 3:1$ $3c$ 1.7:1
21d	2.5, 24	99	3d $2:1$
21e 22	2.5, 24 2.5, 4	98 96	$3e$ 3:1 42.5:1 5 1 4:1
	21c 23	5.0, 24 5.0, 24	85 60

triethylamine (6.8 mL, 50 mmol) in DCM (20 mL) was added 11 (9.5 g, 48 mmol) in DCM (15 mL). The reaction mixture was stirred for 1 h in ice and stirring was continued overnight at room temperature. The resulting solution was extracted with DCM, neutralized with concd HCl, washed with water, $Na₂CO₃$ solution and water again and then dried over anhydrous sodium sulfate. The solvent was then removed in vacuo and remaining precipitate was crystallized from DCM/pet. ether $(40-60)$ to give 5.5 g (42%) of colorless crystals, mp 130–131 °C. MS: $m/z = 269$ (M⁺, 30%). IR: 3326, 3077, 1640, 1598, 1542, 1482, 1457, 1329, 1283, 1232, 1162, 1089, 997, 786, 762, 722. ¹H NMR $(CDCl₃)$: δ 10.38 (s, 1H, NH), 9.61 (s, 1H, OH), 8.33 (dd, 1H, $J=8.0$, 1.5 Hz), 7.55 (dt, 1H, $J=8.3$, 1.5 Hz), 7.19 (t, $2H, J=7.6$ Hz), 7.09 (m, $2H$), 6.98 (d, $1H, J=7.7$ Hz), 6.91 $(t, 1H, J=7.5 Hz)$, 6.21 (m, 1H, CH=), 5.57 (d, 1H, J= 17.2 Hz, CH₂=), 5.50 (d, 1H, $J=10.3$ Hz, CH₂=), 4.80 (d, 2H, $J=6.0$ Hz, OCH₂). Anal. Calcd for C₁₆H₁₅NO₃ (269.3): C 71.36; H 5.61; N 5.20. Found: C 71.10; H 5.61; N 5.44.

3.1.2. Synthesis of 15a–e and 17b,d. General procedure. To a solution of KOH (0.13 g, 2.2 mmol) in methanol (5 mL) was added 12 or 16 (2.2 mmol). The mixture was stirred at room temperature for 15 min and the solvent was

Scheme 4.

Table 3. Catalyst %, yields and E/Z ratios of macrocycles 6–9

Entry	Substrate	(mol%) Catalyst/substrate, reaction time (h)	Yield (%)	Product $E:Z$ ratio
$\mathbf{1}$	30a	2.5, 24	92	6a 3.6:1
$\overline{2}$	30 _b	5, 24	73	6 h 2:1
3	31a	2.5, 24	91	7a3:1
$\overline{4}$	31 _b	5, 24	97	$7b$ 3:1
5	32a	7.5, 3	100	$8a \; 2:1$
6	32 _b	5.3	80	$8b$ 2.6:1
$\overline{7}$	33a	2.5, 4	75	9a 1:1
8	33 _b	2.5, 2	82	$9b$ 2.5:1

then removed in vacuo. To the remaining potassium salt was added DMF (2 mL) and the appropriate dichloro derivatives 14a–e (1 mmol). The reaction mixture was then heated under reflux for 5 min. The mixture was then cooled, diluted with water (20 mL) and the precipitate was collected, washed with cold water and finally crystallized (except 17d, which was obtained as an oil) to give the corresponding dienes 15a–e and 17b, which were also characterized by their R_f values using DCM–pet. ether (40–60)–EtOAc $(6/4/3)$.

3.1.2.1. Compound 15a. Yield 0.8 g (93%); colorless crystals (EtOH), mp 130–131 °C, $R_f=0.7$. LCMS; $m/z = 863$ (M+1). IR: 3400, 3344, 3071, 2933, 2875,

1688, 1663, 1600, 1536, 1482, 1455, 1331, 1293, 1252, 1229, 1163, 1118, 1093, 1047, 999, 915, 750. ¹ H NMR (CDCl₃): δ 10.19 (s, 2H, NH), 8.56 (s, 2H, NH), 8.48 (m, 2H), 8.25 (d, 2H, $J=7.6$ Hz), 8.20 (d, 2H, $J=7.6$ Hz), 7.42 $(t, 2H, J=7.8 \text{ Hz})$, 7.09 $(t, 2H, J=7.4 \text{ Hz})$, 7.02 $(m, 6H)$, 6.71 (m, 6H), 6.42 (d, 2H, $J=7.5$ Hz), 5.83 (m, 2H, CH=), 5.13 (m, 4H, $=CH_2$), 4.44 (s, 4H, OCH₂CO), 4.25 (d, 4H, $J=4.5$ Hz, OCH₂CH= $)$, 3.19 (s, 4H, OCH₂). Anal. Calcd for $C_{50}H_{46}N_4O_{10}$ (862.9): C 69.59; H 5.37; N 6.49. Found: C 69.50; H 5.52; N 6.77.

3.1.2.2. Compound 15b. Yield 0.8 g (91%); colorless crystals (EtOH), mp 130–131 °C, R_f =0.6. LCMS; m/z = 877 (M+1). IR: 3400, 3344, 3064, 2937, 1674, 1601, 1536, 1484, 1456, 1335, 1293, 1227, 1120, 1051, 998, 931, 851, 752. ¹H NMR (CDCl₃): δ 10.52 (s, 2H, NH), 8.70 (dd, 2H, $J=7.8$, 1.7 Hz), 8.63 (s, 2H, NH), 8.29 (d, 4H, $J=7.8$ Hz), 7.51 (dt, 2H, $J=7.7$, 1.8 Hz), 7.09 (m, 6H), 6.95 (d, 2H, $J=$ 8.3 Hz), 6.86 (m, 4H), 6.67 (dt, 2H, $J=7.8$, 1.4 Hz), 6.34 (d, 2H, $J=8.0$ Hz), 5.98 (m, 2H, CH=), 5.30 (d, 2H, $J=$ 17.2 Hz, $=CH_2$), 5.19 (d, 2H, $J=10.5$ Hz, $=CH_2$), 4.69 (s, 4H, OCH₂CO), 4.57 (d, 4H, $J=5.7$ Hz, OCH₂CH=), 3.43 (t, 4H, $J=5.2$ Hz, OCH₂CH₂), 1.18 (quint, 2H, $J=5.2$ Hz, OCH₂CH₂). Anal. Calcd for C₅₁H₄₈N₄O₁₀ (876.9): C 69.85; H 5.52; N 6.39. Found: C 69.63; H 5.43; N 6.29.

3.1.2.3. Compound 15c. Yield 0.7 g (77%); colorless crystals (EtOH), mp 160–161 °C, R_f =0.6. LCMS; m/z = 907 (M+1). IR: 3391, 3341, 3069, 2936, 2874, 1691, 1658, 1601, 1535, 1483, 1455, 1333, 1295, 1254, 1229, 1135, 1118, 1047, 753. ¹H NMR (CDCl₃): δ 10.49 (s, 2H, NH), 8.67 (d, 2H, $J=8.0$ Hz), 8.63 (s, 2H, NH), 8.36 (d, 2H, $J=$ 8.0 Hz), 8.27 (d, 2H, $J=8.0$ Hz), 7.48 (t, 2H, $J=8.0$ Hz), 7.13 (t, 2H, $J=8.0$ Hz), 7.05 (m, 4H), 6.92 (m, 8H), 6.41 (d, 2H, $J=8.0$ Hz), 5.99 (m, 2H, CH=), 5.29 (d, 2H, $J=$ 16.5 Hz, $=CH_2$), 5.19 (d, 2H, $J=10.4$ Hz, $=CH_2$), 4.64 (s, 4H, OCH₂CO), 4.53 (d, 4H, $J=5.6$ Hz, OCH₂CH=), 3.43 (t, 4H, $J=4.4$ Hz, OCH₂CH₂O), 3.95 (t, 4H, $J=4.4$ Hz, OCH₂CH₂O). Anal. Calcd for $C_{52}H_{50}N_4O_{11}$ (907.0): C 68.86; H 5.56; N 6.18. Found: C 68.71; H 5.45; N 6.47.

3.1.2.4. Compound 15d. Yield 0.6 g (63%); colorless crystals (EtOH/CHCl₃), mp 143–144 °C, R_f =0.4. LCMS; $m/z = 952$ (M + 1). IR: 3396, 3344, 3064, 3012, 2920, 1663, 1600, 1535, 1483, 1456, 1333, 1293, 1254, 1226, 1117, 1048, 932, 752. ¹H NMR (CDCl₃): δ 10.49 (s, 2H, NH), 8.83 $(s, 2H, NH)$, 8.67 (dd, 2H, $J=7.6$, 1.9 Hz), 8.40 (dd, 2H, $J=$ 7.7, 1.6 Hz), 8.29 (dd, 2H, $J=7.8$, 1.6 Hz), 7.45 (dt, 2H, $J=$ 8.0, 1.6 Hz), 7.10 (t, 2H, $J=7.6$ Hz), 7.02 (m, 8H), 6.92 (m, 4H), 6.76 (dd, 2H, $J=8.0$, 1.2 Hz), 6.05 (m, 2H, CH=), 5.33 (d, 2H, $J=17.2$ Hz, $=CH_2$), 5.24 (d, 2H, $J=10.5$ Hz, $=CH₂$), 4.72 (s, 4H, OCH₂CO), 4.57 (d, 4H, J = 5.6 Hz, OCH₂CH=), 3.67 (t, 4H, $J=5.2$ Hz, OCH₂CH₂O), 3.15 (t, 4H, $J=5.2$ Hz, OCH₂CH₂O), 3.07 (s, 4H, OCH₂CH₂O). Anal. Calcd for C₅₄H₅₄N₄O₁₂ (951.0): C 68.20; H 5.72; N 5.89. Found: C 67.98; H 5.57; N 6.10.

3.1.2.5. Compound 15e. Yield 0.8 g (85%); colorless crystals (EtOH), mp 228–229 °C, R_f =0.8. LCMS; m/z = 939 (M+1). IR: 3393, 3322, 3070, 3005, 2935, 2884, 1691, 1654, 1599, 1527, 1482, 1454, 1332, 1290, 1248, 1231, 1188, 1136, 1114, 1091, 1044, 997, 949, 929, 895, 790, 753. ¹H NMR (CDCl₃): δ 10.20 (s, 2H, NH), 8.78 (d, 2H, J= 7.7 Hz), 8.62 (s, 2H, NH), 8.43 (m, 2H), 8.23 (d, 2H, $J=$ 7.5 Hz), 7.45 (t, 2H, $J=7.5$ Hz), 6.96 (m, 16H), 6.43 (m, 2H), 6.38 (m, 2H), 5.73 (m, 2H, CH=), 5.20 (d, 2H, $J=$ 17 Hz, $=CH_2$), 5.08 (d, 2H, $J=10.2$ Hz, $=CH_2$), 4.62 (s, 4H, OCH₂CO), 4.36 (s, 4H, OCH₂Ar), 4.27 (d, 4H, $J=$ 5.5 Hz, OCH₂CH=). Anal. Calcd for C₅₆H₅₀N₄O₁₀ (939): C 71.63; H 5.37; N 5.97. Found: C 71.34; H 5.46; N 6.20.

3.1.2.6. Compound 17b. Yield 0.4 g (53%); colorless crystals (EtOH), mp 126–127 °C, R_f =0.7. LCMS; m/z = 753 (M+1). IR: 3388, 3270, 3070, 2934, 2879, 1679, 1603, 1537, 1486, 1456, 1290, 1260, 1119, 750. ¹ H NMR (CDCl₃): δ 9.03 (s, 2H, NH), 8.63 (s, 2H, NH), 8.44 (m, 2H), 8.30 (m, 2H), 7.06 (m, 4H), 6.97 (m, 4H), 6.92 (m, 2H), 6.58 (m, 2H), 5.82 (m, 2H, CH=), 5.27 (d, 2H, $J=17.2$ Hz, $=CH_2$), 5.15 (d, 2H, $J=10.4$ Hz, $=CH_2$), 4.74 (s, 4H, OCH₂CO), 4.06 (t, 4H, $J=5.6$ Hz, OCH₂CH=), 4.05 (s, 4H, OCH₂CO), 3.71 (t, 4H, J=5.5 Hz, OCH₂CH₂), 1.72 (quint, 2H, $J=5.5$ Hz, OCH₂CH₂). Anal. Calcd for $C_{41}H_{44}N_{4}O_{10}$ (752.8): C 65.41; H 5.89; N 7.44. Found: C 65.32; H, 5.85; N 7.60.

3.1.2.7. Compound 17d. Yield 0.12 g (15%); pale yellow oil purified using column chromatography using EtOAc/pet. ether (40–60) as an eluent, $R_f = 0.6/EtOAc$. LCMS; $m/z = 827$ (M+1). IR: 3389, 3070, 2923, 2854,

1689, 1603, 1535, 1486, 1456, 1292, 1255, 1207, 1117, 1045, 930, 750. ¹H NMR (CDCl₃): δ 9.06 (s, 2H, NH), 8.76 $(s, 2H, NH)$, 8.43 (m, 2H), 8.34 (d, 2H, $J=7.9$ Hz), 6.92 (m, 12H), 5.80 (m, 2H, CH=), 5.25 (dd, 2H, $J=17.3$, 1.0 Hz, $=CH_2$), 5.11 (d, 2H, $J=10.2$ Hz, $=CH_2$), 4.71 (s, 4H, OCH₂CO), 4.08 (s, 4H, OCH₂CO), 4.04 (d, 4H, $J=5.6$ Hz, OCH₂CH=), 3.99 (t, 4H, $J=4.8$ Hz, OCH₂CH₂O), 3.49 (t, 4H, $J=4.8$ Hz, OCH₂CH₂O), 3.33 (s, 4H, OCH₂CH₂O). Anal. Calcd for $C_{44}H_{50}N_4O_{12}$ (826.9): C 63.91; H 6.09; N 6.78. Found: C 63.68; H 6.31; N 6.77.

3.1.3. Methyl 2-(2-allyloxyethoxy)benzoate. A mixture of methyl salicylate (10 g, 65 mmol), 2-allyloxyethyl p-toluenesulfate (8.5 g, 33 mmol) and anhydrous K_2CO_3 (4.5 g, 33 mmol) in dry acetone (50 mL) was heated under reflux for 48 h. The solid precipitate was filtered off and the solvent was evaporated in vacuo. The remaining oil was dissolved in DCM, washed twice with aq KOH (10%), dried over anhydrous $Na₂SO₄$ and the solvent was then removed in vacuo and the remaining oil was purified by column chromatography over silica gel using DCM/pet ether (40–60) as an eluent to give 6.8 g (87%) of colorless oil, $R_f = 0.3$ [DCM/pet. ether (40–60) 2:1]. LCMS; $m/z = 251$ $(M+1)$. IR: 3078, 2947, 2871, 1727, 1600, 1491, 1450, 1304, 1252, 1133, 1088, 1050, 926, 758. ¹H NMR (CDCl₃): δ 7.80 (dd, 1H, $J=8.0$, 1.6 Hz), 7.46 (dt, 1H, $J=8.8$, 1.7 Hz), 7.01 (m, $2H$), 5.97 (m, $1H$, CH=), 5.33 (dd, $1H$, $J=17.2$, 1.6 Hz, CH₂=), 5.22 (dd, 1H, $J=9.4$, 0.9 Hz, CH₂=), 4.23 (t, 2H, $J=5.08$ Hz, CH₂), 4.15 (d, 2H, $J=$ 5.5 Hz, CH2), 3.89 (m, 4H, CH2), 3.89 (s, 3H, CH3). Anal. Calcd for $C_{13}H_{16}O_4$ (236.3): C 66.09; H 6.83. Found: C 65.95; H 6.62.

3.1.4. 2-(2-Allyloxyethoxy)benzoic acid. A mixture of methyl 2-(2-allyloxyethoxy)benzoate (1.2 g, 5 mmol) in aq KOH (10%, 10 mL) was heated under reflux for 24 h. The mixture was neutralized with concd HCl and extracted with DCM, washed with water, separated and dried over anhydrous sodium sulfate. The solvent was then removed in vacuo to give 0.8 g (72%) of colorless oil, which was used in the next step without further purification. LCMS; $m/z=$ 223 (M + 1). IR: 3300–2871 (br) 3071, 2928, 2871, 1731, 1603, 1486, 1456, 1387, 1351, 1297, 1240, 1162, 1124, 1038, 926, 756. ¹H NMR (CDCl₃): δ 9.89 (br, 1H), 8.17 (dd, 1H, $J=7.8$, 1.1 Hz), 7.56 (dt, 1H, $J=7.9$, 1.6 Hz), 7.15 (t, 1H, $J=7.4$ Hz), 7.06 (d, 1H, $J=8.4$ Hz), 5.93 (m, 1H, CH=), 5.32 (dd, 1H, $J=16$, 1.1 Hz, CH₂=), 5.24 (d, 1H, $J=10.4$ Hz, CH₂=), 4.39 (t, 2H, $J=4.5$ Hz, CH₂), 4.09 (d, 2H, $J=5.8$ Hz, CH₂), 3.87 (t, 2H, $J=4.5$ Hz, CH₂).

3.1.5. 2-(2-Allyloxyethoxy)benzoyl chloride 19. A mixture of 2-(2-allyloxyethoxy)benzoic acid (2.22 g, 10 mmol) and SOC_{2} (1.5 mL, 20 mmol) was heated under reflux at 100 °C for 1.5 h. The excess $S OCl₂$ was removed in vacuo to give 2.2 g (92%) of colorless oil, which was used directly in the next step without further purification.

3.1.6. Preparation of the bis-amides 21–23. General **procedure.** A. To a cold solution $(0-5 \degree C)$ of each of 20b–e, ethylenediamine or o-phenylenediamine (0.8 mmol) and triethylamine (0.25 g, 2.5 mmol) in DCM (20 mL), was added 19 (0.44 g, 1.7 mmol) in DCM (5 mL) dropwise with stirring. After complete addition the reaction mixture was

stirred for 1 h at $0-5$ °C and stirring was continued overnight at room temperature. The organic layer was extracted with DCM, washed with concd HCl (5 mL), water twice, $NaHCO₃$ solution and finally water. The organic layer was separated, dried over anhydrous sodium sulfate, the solvent was then removed in vacuo and the remaining material was purified by column chromatography. All compounds were characterized by R_f values using the solvent mixture DCM–pet. ether (40–60)–EtOAc in (6/4/2).

B. To a mixture of each of 1,2-bis(2-hydroxybenzamido) ethane and 1,2-bis(2-hydroxybenzamido)benzene (0.8 mmol), anhydrous K_2CO_3 (0.4 g, 3 mmol) and 2-allyloxyethyl p-toluenesulfonate (0.5 g, 2 mmol) in dry DMF (2 mL) was heated at 100 \degree C for 10 h. The product was then extracted with DCM, washed several times with water, separated and dried over anhydrous sodium sulfate. The solvent was then removed in vacuo and the remaining product was purified to give the corresponding products 22 and 23, respectively.

3.1.6.1. Compound 21b. Yield 0.26 g (50%), colorless oil, $R_f=0.8$. LCMS; $m/z=667$ (M+1). IR: 3346, 3072, 2938, 2878, 1660, 1599, 1535, 1481, 1454, 1332, 1291, 1252, 1234, 1162, 1134, 1117, 1091, 1048, 995, 927, 753. ¹H NMR (CDCl₃): δ 10.22 (s, 2H, NH), 8.46 (d, 2H, J= 7.6 Hz), 8.27 (dd, 2H, $J=8.0$, 1.8 Hz), 7.45 (dt, 2H, $J=7.9$, 1.7 Hz), 7.14 (t, 2H, $J=7.5$ Hz), 6.98 (m, 6H), 6.87 (dd, 2H, $J=8.0$, 1.4 Hz), 5.67 (m, 2H, CH=), 5.06 (dd, 2H, $J=17.3$, 1.5 Hz, CH₂=), 4.99 (dd, 2H, $J=10.4$, 1.2 Hz, CH₂=), 4.30 (m, 8H, CH₂), 3.86 (d, 4H, $J=5.6$ Hz), 3.72 (t, 4H, $J=$ 4.8 Hz), 2.33 (quint, 2H, $J=4.6$ Hz). ¹³C/DEPT NMR (CDCl3): d 163.1 (C), 156.4 (C), 147.9 (C), 133.9 (CH), 132.9 (CH), 132.1 (CH), 128.2 (C), 123.9 (CH), 122.7 (C), 121.8 (CH), 121.7 (CH), 121.2 (CH), 117.1 (CH₂), 113.9 (CH), 111.6 (CH), 72.0 (CH₂), 69.2 (CH₂), 67.7 (CH₂), 65.0 (CH₂), 28.7 (CH₂). Anal. Calcd for C₃₉H₄₂N₂O₈ (666.8): C 70.25; H 6.35; N 4.20. Found: C 69.89; H 6.33; N 4.09.

3.1.6.2. Compound 21c. Yield 0.33 g (59%), colorless oil, $R_f=0.7$. LCMS; $m/z=697$ (M+1). IR: 3342, 3072, 2932, 2874, 1660, 1599, 1534, 1481, 1454, 1332, 1291, 1235, 1132, 1093, 1048, 929, 753. ¹H NMR (CDCl₃): δ 10.24 (s, 2H, NH), 8.53 (m, 2H), 8.22 (dd, 2H, $J=7.6$, 1.6 Hz), 7.40 (dt, 2H, $J=7.8$, 2 Hz), 7.10 (t, 2H, $J=7.6$ Hz), 7.04 (m, 4H), 6.95 (d, 2H, $J=8.4$ Hz), 6.86 (m, 2H), 5.72 $(m, 2H, CH=), 5.11$ (dd, $2H, J=17.3, 1.6$ Hz, $CH_2=), 5.03$ (dd, 2H, $J=10.5$, 1 Hz, CH₂=), 4.31 (t, 4H, $J=4.9$ Hz, OCH₂CH₂), 4.16 (t, 4H, $J=4.7$ Hz, OCH₂CH₂), 3.90 (dd, 4H, $J=5.7$, 1 Hz, OCH₂CH=), 3.84 (t, 4H, $J=4.7$ Hz, OCH₂CH₂), 3.78 (t, 4H, $J=4.9$ Hz, OCH₂CH₂). Anal. Calcd for $C_{40}H_{44}N_2O_9$ (696.8): C 68.95; H 6.36; N 4.02. Found: C 69.15; H 6.18; N 3.98.

3.1.6.3. Compound 21d. Yield 0.26 g (44%), colorless crystals, mp 56–57 °C, R_f =0.5. LCMS; m/z =741 (M+1). IR: 3340, 3073, 2927, 2874, 1660, 1599, 1536, 1480, 1456, 1333, 1291, 1254, 1235, 1221, 1162, 1134, 1115, 1091, 1048, 997, 928, 754. ¹H NMR (CDCl₃): δ 10.24 (s, 2H, NH), 8.54 (dd, 2H, $J=7.4$, 2.3 Hz), 8.24 (dd, 2H, $J=7.8$, 1.4 Hz), 7.43 (dt, 2H, $J=8.4$, 1.5 Hz), 7.12 (t, 2H, $J=7.5$ Hz), 7.05 $(m, 6H), 6.92$ (dd, 2H, $J=7.7, 2.3$ Hz), 5.76 $(m, 2H, CH=)$, 5.14 (dd, 2H, $J=17.2$, 1.3 Hz, CH₂=), 5.06 (d, 2H, $J=10.3$ Hz, CH₂=), 4.38 (t, 4H, $J=4.9$ Hz, CH₂), 4.19 (t, 4H, $J=4.7$ Hz, OCH₂), 3.95 (d, 4H, $J=5.6$ Hz, OCH₂CH = \leq 3.83 (t, 4H, J = 4.9 Hz, OCH₂), 3.75 (t, 4H, $J=4.7$ Hz, OCH₂), 3.52 (s, 4H, CH₂). ¹³C NMR (CDCl₃): δ 163.3, 156.6, 148.0, 134.2, 132.9, 132.3, 128.8, 124.8, 123.9, 121.8, 121.6, 121.5, 117.2, 113.6, 112.3, 72.2, 70.8, 69.4, 69.1, 68.5, 68.0. Anal. Calcd for $C_{42}H_{48}N_2O_{10}$ (740.9): C 68.09; H 6.53; N 3.78. Found: C 67.87; H 6.65; N 3.86.

3.1.6.4. Compound 21e. Yield 0.27 g (46%), colorless oil, R_f =0.9. LCMS; m/z =729 (M+1). IR: 3343, 3072, 2933, 2874, 1661, 1599, 1535, 1480, 1454, 1332, 1293, 1233, 1162, 1133, 1092, 1046, 100, 922, 752. ¹ H NMR $(CDCl₃)$: δ 10.40 (s, 2H, NH), 8.54 (d, 2H, $J=8.0$ Hz), 8.30 $(dd, 2H, J=7.8, 1.2 Hz$), 7.53 (m, 2H), 7.45 (dt, 2H, $J=7.7$, 1.4 Hz), 7.39 (m, 2H), 7.13 (t, 2H, $J=7.5$ Hz), 6.98 (m, 4H), 6.87 (m, 4H), 5.63 (m, 2H, CH=), 5.29 (s, 4H, OCH₂Ar), 5.04 (dd, 2H, $J=17.5$, 1 Hz, CH₂=), 4.98 (d, 2H, $J=$ 10.3 Hz, CH₂=), 3.87 (t, 4H, $J=4.7$ Hz, OCH₂), 3.76 (d, 4H, $J=5.4$ Hz, OCH₂), 3.37 (t, 4H, $J=4.7$ Hz, OCH₂). Anal. Calcd for $C_{44}H_{44}N_2O_8$ (728.9): C 72.51; H 6.09; N 3.84. Found: C 72.56; H 5.99; N 4.10.

3.1.6.5. Compound 22. Yield 0.19 g (51%, A), 0.2 g (52%, B), colorless crystals from EtOAc, mp $95-96$ °C. LCMS; $m/z = 469$ (M+1). IR: 3354, 3071, 2943, 2861, 1646, 1600, 1539, 1483, 1448, 1296, 1245, 1087, 1051, 1020, 999, 925, 766. ¹H NMR (CDCl₃): δ 8.46 (s, 2H, NH), 8.19 (dd, 2H, $J=7.7$, 1.3 Hz), 7.42 (dt, 2H, $J=7.7$, 1.3 Hz), 7.08 (t, 2H, $J=7.5$ Hz), 6.96 (d, 2H, $J=8.2$ Hz), 5.85 (m, 2H, CH=), 5.27 (d, 2H, $J=17.7$ Hz, CH₂=), 5.20 (d, 2H, $J=13.8$ Hz, CH₂ $=$), 4.23 (t, 4H, $J=4.6$ Hz, OCH₂CH₂), 3.97 (d, 4H, $J=5.6$ Hz, OCH₂CH=), 3.73 (t, 4H, $J=$ 4.6 Hz, OCH₂CH₂), 3.68 (m, 4H, NCH₂). Anal. Calcd for $C_{26}H_{32}N_{2}O_{6}$ (468.5): C 66.65; H 6.88; N 5.98. Found: C 66.52; H 6.84; N 6.11.

3.1.6.6. Compound 23. Yield 0.88 g (85%, A), 0.74 g (71%, B), colorless oil, $R_f = 0.8$. LCMS; $m/z = 517$ (M+1). IR: 3320, 3075, 2934, 2869, 1662, 1598, 1529, 1510, 1480, 1454, 1300, 1231, 1111, 1045, 927, 756. ¹H NMR (CDCl₃): δ 10.02 (s, 2H, NH), 8.22 (dd, 2H, J=7.8, 1.4 Hz), 7.80 (m, 2H), 7.46 (dt, 2H, $J=8.0$, 1.5 Hz), 7.26 (m, 2H), 7.10 (t, 2H, $J=7.5$ Hz), 7.03 (d, 2H, $J=8.2$ Hz), 5.68 (m, 2H, CH=), 5.11 (dd, 2H, $J=17.3$, 1.4 Hz, CH₂=), 5.05 (d, 2H, $J=$ 10.3 Hz, CH₂=), 4.22 (t, 4H, $J=4.8$ Hz, OCH₂CH₂), 3.86 (d, 4H, $J=5.6$ Hz, OCH₂CH=), 3.65 (t, 4H, $J=4.8$ Hz, OCH₂CH₂). Anal. Calcd for C₃₀H₃₂N₂O₆ (516.6): C 69.75; H 6.24; N 5.42. Found: C 69.50; H 6.28; N 5.70.

3.1.7. 2-(2-Allyloxyethoxy)acetanilide. A. A mixture of o-acetamidophenol (1.5 g, 10 mmol), 2-allyloxyethyl p-toluenesulfate (2.56 g, 10 mmol) and anhydrous K_2CO_3 (5.5 g) in dry DMF (5 mL) was heated at 100 °C for 4 h. Crushed ice was then added to the mixture, the precipitate was washed three times with cold KOH soln (10%) and then with cold water twice and then extracted with DCM, and dried over anhydrous $Na₂SO₄$. The solvent was then removed in vacuo to give 1.98 g (86%) of the product as a pale yellow oil, which was used in the next step without further purification. MS: $m/z = 235.1$ (M⁺, 70%). IR: 3531, 3423, 3309, 2932, 2872, 1684, 1601, 1531, 1487, 1450, 1371, 1327, 1289, 1255, 1213, 1118, 1036, 929, 752.

¹H NMR (CDCl₃): δ 8.27 (dd, 1H, J=7.6, 1.5 Hz), 8.24 (s, 1H, NH), 6.90 (m, 2H), 6.82 (dd, 1H, J=7.6, 1.5 Hz), 5.83 (m, 1H, $=$ CH), 5.23 (dd, 1H, $J=17.1$, 1.2 Hz, $=$ CH₂), 5.13 (d, 1H, $J=10.3$ Hz, $=CH_2$), 4.04 (t, 2H, $J=4.5$ Hz, OCH₂CH₂), 3.97 (d, 2H, $J=5.6$ Hz, CH₂CH=), 3.66 (t, 2H, $J=4.5$ Hz, OCH₂CH₂), 2.07 (s, 3H, CH₃).

B. To a solution of KOH (0.56 g, 10 mmol) in methanol (15 mL) , was added *o*-acetamidophenol $(1.5 \text{ g}, 10 \text{ mmol})$. The mixture was then stirred at room temperature for 15 min and the solvent was then removed in vacuo. To the remaining potassium salt was added DMF (2 mL) and 2-allyloxyethyl p -toluenesulfate (2.56 g, 10 mmol). The reaction mixture was then heated under reflux for 5 min. The mixture was cooled, diluted with water (20 mL) and extracted with DCM, washed with cold water, dried over $Na₂SO₄$ and evaporated to give 2.09 g (91%) of the product as a pale yellow oil, which was used in the next step without further purification.

3.1.8. 2-(2-Allyloxyethoxy)aniline hydrochloride 24b. A mixture of 2-(2-allyloxyethoxy)acetanilide (10 mmol) and concd HCl (2 mL) in ethanol (10 mL) was heated under reflux for 2 h. The solvent was then removed in vacuo and the residue obtained was washed with ether and crystallized from ethyl acetate to give 1.5 g (65%) of the product as colorless crystals, mp 124 °C. MS: $m/z = 193$ [(C₁₁H₁₅NO₂)⁺, 30%]. IR: 3412, 2873, 2591, 1630, 102, 1461, 1267, 1098, 1026, 931, 755. ¹H NMR (CDCl₃): δ 10.49 (br s, 3H, NH₂HCl), 7.74 (d, 1H, $J=7.5$ Hz), 7.33 (t, 1H, $J=7.5$ Hz), 7.00 (m, 2H), 5.84 (m, 1H, CH=), 5.21 (d, 1H, $J=17.2$ Hz, CH₂=), 5.09 (d, 1H, $J=10.2$ Hz, CH₂ $=$), 4.25 (br, 2H, OCH₂CH₂), 4.03 (d, 2H, $J=5$ Hz, CH₂CH=), 3.92 (br, 2H, OCH₂CH₂). Anal. Calcd for $C_{11}H_{16}NO_2Cl$ (229.7): C 57.52; H 7.02; N 6.10. Found: C 57.26; H 6.90; N 6.28.

3.1.9. Chloroacetanilides 25a,b. General procedure. To a cold $(0-5^{\circ}C)$ solution of each of 24a,b (50 mmol) and triethylamine (10.6 mL, 76 mmol) in DCM (25 mL) was added dropwise with stirring chloroacetyl chloride (3.8 mL, 50 mmol) in DCM (25 mL). The reaction mixture was stirred for 1 h in ice and stirring was continued overnight at room temperature. The resulting solution was neutralized with HCl, washed with $Na₂CO₃$ solution and water, extracted with DCM, dried over sodium sulfate and the solvent was removed in vacuo to give the corresponding product 25a,b.

3.1.9.1. o-Allyloxychloroacetanilide 25a. Yield 10.8 g (96%), pale yellow oil. MS: $m/z = 225$ (M⁺, 20%). IR: 3388, 2872, 1683, 1603, 1537, 1487, 1456, 1411, 1337, 1291, 1255, 1207, 1117, 997, 914, 748. ¹H NMR (CDCl₃): δ 9.06 (s, 1H, NH), 8.35 (dd, 1H, $J=8$, 1.2 Hz), 7.07 (dt, 1H, $J=8$, 1.2 Hz), 6.98 (t, 1H, $J=8$ Hz), 6.88 (d, 1H, $J=8$ Hz), 6.06 (m, 1H, CH=), 5.47 (dd, 1H, $J=16$, 1.1 Hz, CH₂=), 5.32 (dd, 1H, $J=9.4$, 1 Hz, CH₂ $=$), 4.58 (d, 2H, $J=4.9$ Hz, $CH₂CH=$), 4.20 (s, 2H, COCH₂). Anal. Calcd for $C_{11}H_{12}NO_2Cl$ (225.7): C 58.55; H 5.36; N 6.21. Found: C 58.32; H 5.37; N 6.40.

3.1.9.2. N-[2-(2-Allyloxyethoxy)phenyl]chloroacetamide 25b. Yield 11.3 g (84%), pale yellow oil. MS: $m/z = 271$ (M + 1, 25%), 269 (M⁺, 78%). IR: 3505, 3383, 3075, 3014, 2938, 2871, 1763, 1689, 1603, 1534, 1487, 1454, 1408, 1338, 1290, 1257, 1210, 1118, 1043, 996, 928, 752. ¹H NMR (CDCl₃): δ 9.11 (s, 1H, NH), 8.34 (d, 1H, J= 7.8 Hz), 7.09 (t, 1H, $J=7.8$ Hz), 7.01 (t, 1H, $J=7.8$ Hz), 6.92 (d, 1H, $J=7.8$ Hz), 5.96 (m, 1H, CH=), 5.32 (d, 1H, $J=17.3$ Hz, CH₂=), 5.23 (d, 1H, $J=10.4$ Hz, CH₂=), 4.23 $(t, 2H, J=4.5 Hz, OCH₂CH₂), 4.20 (s, 2H, COCH₂), 4.10$ (d, 2H, $J=5.6$ Hz, CH₂CH=), 3.83 (t, 2H, $J=4.5$ Hz, OCH₂CH₂). ¹³C NMR (CDCl₃): δ 163.6, 147.5, 133.3, 127.2, 124.5, 121.5, 119.7, 117.5, 112, 72.4, 68.6, 68.4, 43.1. Anal. Calcd for $C_{13}H_{16}NClO_3$ (269.7): C 57.89; H 5.98; N 5.19. Found: C 57.62; H 5.75; N 5.40.

3.1.10. Synthesis of compounds 30–33. General procedures. A mixture of each of the appropriate chloroacetanilide 25a,b (1 mmol) and the appropriate phenolic derivative $26-29$ (1 mmol) and powdered K_2CO_3 (4 mmol) in anhydrous DMF (1.5 mL) was stirred at 100 °C for 1 h. After cooling crushed ice was added and the mixture was extracted with DCM. The organic layer was separated, washed with water and dried over anhydrous sodium sulfate. The solvent was removed in vacuo and remaining product was crystallized from the proper solvent.

3.1.10.1. Compound 30a. Yield 0.24 g (49%), colorless crystals (EtOH), mp 144 °C. LCMS; $m/z = 489$ (M + 1). IR: 3436, 3383, 3023, 2970, 2868, 1684, 1601, 1539, 1503, 1485, 1456, 1421, 1336, 1292, 1253, 1203, 1131, 1115, 1046, 1017, 928, 751. ¹H NMR (CDCl₃): δ 9.19 (s, 2H, NH), 8.43 (dd, 2H, $J=7.5$, 1.9 Hz), 7.03 (m, 8H), 6.59 (dd, 2H, $J=7.8$, 1.5 Hz), 5.86 (m, 2H, =CH), 5.30 (dd, 2H, $J=17.3$, 1.1 Hz, = CH₂), 5.16 (dd, 2H, $J = 10.6$, 0.9 Hz, = CH₂), 4.70 (s, 4H, OCH₂CO), 4.15 (d, 4H, $J=5.0$ Hz, OCH₂CH=). Anal. Calcd for $C_{28}H_{28}N_2O_6$ (488.6): C 68.84; H 5.78; N 5.73. Found: C 68.62; H 5.88; N 5.76.

3.1.10.2. Compound 30b. Yield 0.5 g (87%), pale yellow. MS: $m/z = 576.2$ (M⁺, 100%). IR: 3393, 3070, 2932, 2863, 1690, 1601, 1536, 1503, 1486, 1457, 1256, 1211, 1118, 1048, 928, 750. ¹H NMR (CDCl₃): δ 9.18 (s, 2H, NH), 8.39 (dd, 2H, $J=7.8$, 1.6 Hz), 7.01 (m, 8H), 6.67 (d, 2H, $J=7.9$ Hz), 5.74 (m, 2H, $=CH$), 5.14 (dd, 2H, $J=$ 17.2, 1.4 Hz, $=CH_2$), 5.02 (d, 2H, $J=10.4$ Hz, $=CH_2$), 4.72 (s, 4H, OCH₂CO), 3.91 (d, 4H, $J=5.5$ Hz, OCH₂CH = \leq 3.83 (t, 4H, J = 4.7 Hz, OCH₂CH₂O), 3.63 (t, 4H, $J=4.7$ Hz, OCH₂CH₂O). Anal. Calcd for $C_{32}H_{36}N_2O_8$ (576.7): C 66.65; H 6.29; N 4.86. Found: C 64.18; H 5.93; N 5.86.

3.1.10.3. Compound 31a. Yield 0.12 g (18%), colorless crystals (acetone), mp 141-142 °C. LCMS; $m/z = 665$ (M+1). IR: 3387, 3061, 2918, 2852, 1690, 1599, 1534, 1507, 1483, 1457, 1360, 1326, 1272, 1253, 1210, 1149, 1121, 1066, 992, 924, 813, 751. ¹H NMR (CDCl₃): δ 8.42 (s, 2H, NH), 8.08 (dd, 2H, $J=7.9$, 1.4 Hz), 7.97 (d, 2H, $J=9.0$ Hz), 7.87 (d, 2H, $J=8.2$ Hz), 7.43 (d, 2H, $J=9$ Hz), 7.37 (t, 2H, $J=7.4$ Hz), 7.28 (t, 2H, $J=9$ Hz), 7.18 (d, 2H, $J=8.4$ Hz), 6.98 (t, 2H, $J=7.8$ Hz), 6.89 (d, 2H, $J=7.6$ Hz), 6.72 (d, 2H, $J=7.8$ Hz), 5.57 (m, 2H, CH=), 5.07 (dd, 2H, $J=17.3$, 1.3 Hz, $=CH_2$), 4.95 (dd, 2H, $J=10.6$, 1.2 Hz, $=CH_2$), 4.59, 4.52 (2d, 4H, $J=15.5$ Hz, OCH₂CO), 4.25 (d, 4H, $J=$ 5 Hz, OCH₂CH=). Anal. Calcd for $C_{42}H_{36}N_2O_6$ (664.8): C 75.89; H 5.46; N 4.21. Found: C 75.67; H 5.62; N 4.22.

3.1.10.4. Compound 31b. Yield 0.7 g (93%), colorless crystals (EtOH), mp 144–145 °C. LCMS; $m/z=753$ $(M+1)$. IR: 3391, 3063, 2870, 1693, 1601, 1533, 1484, 1457, 1325, 1254, 1211, 1147, 1114, 1063, 927, 754. ¹H NMR (CDCl₃): δ 8.47 (s, 2H, NH), 8.07 (dd, 2H, $J=7.9$, 1.2 Hz), 7.97 (d, 2H, $J=9$ Hz), 7.88 (d, 2H, $J=8.2$ Hz), 7.44 (d, 2H, $J=9$ Hz), 7.39 (t, 2H, $J=7.6$ Hz), 7.28 (t, 2H, $J=7.6$ Hz), 7.20 (d, 2H, $J=8.4$ Hz), 7.03 (dt, 2H, $J=7.8$, 1.2 Hz), 6.92 (t, 2H, $J=7.7$ Hz), 6.81 (d, 2H, $J=8.0$ Hz), 5.70 (m, 2H, CH=), 5.09 (dd, 2H, $J=17.4$, 1.6 Hz, CH₂=), 5.04 (d, 2H, $J=10.5$ Hz, CH₂ $=$), 4.55 (s, 4H, OCH₂CO), 3.89 (m, 4H, OCH₂CH₂), 3.67 (d, 4H, $J=5.5$ Hz, $OCH_2CH=$), 3.22 (m, 4H, OCH_2CH_2). Anal. Calcd for $C_{46}H_{44}N_2O_8$ (752.9): C 73.39; H 5.89; N 3.72. Found: C 73.07; H 5.96; N 3.91.

3.1.10.5. Compound 32a. Yield 0.2 g (29%), colorless crystals (EtOH), mp $167-168$ °C. LCMS: $m/z=679$ $(M+1)$. IR: 3397, 3374, 3350, 1698, 1640, 1602, 1542, 1489, 1453, 1299, 1254, 1223, 1116, 1043, 1005, 750. ¹H NMR (CDCl₃): δ 8.65 (s, 2H, NH), 8.53 (s, 2H, NH), 8.25 $(d, 2H, J=7.8 \text{ Hz})$, 8.11 $(d, 2H, J=7.4 \text{ Hz})$, 7.35 $(t, 2H, J=$ 7.5 Hz), 7.03 (m, 4H), 6.92 (t, 2H, $J=7.7$ Hz), 6.83 (d, 2H, $J=8.2$ Hz), 6.78 (d, 2H, $J=8.0$ Hz), 5.81 (m, 2H, CH=), 5.19 (m, 4H, CH₂=), 4.72 (s, 4H, OCH₂CO), 4.41 (d, 4H, $J=4.7$ Hz, OCH₂CH= $)$, 3.82 (s, 4H, NCH₂). Anal. Calcd for $C_{38}H_{38}N_4O_8$ (678.8): C 67.24; H 5.64; N 8.25. Found: C 66.98; H 5.71; N 8.46.

3.1.10.6. Compound 32b. Yield 0.65 g (85%), colorless crystals (EtOAc), mp 67–68 °C. LCMS; $m/z = 767$ (M + 1). IR: 3385, 3334, 2929, 2862, 1692, 1642, 1601, 1537, 1485, 1453, 1295, 1255, 1206, 1161, 1117, 1046, 928, 752. ¹H NMR (CDCl₃): δ 8.86 (s, 4H, NH), 8.14 (d, 2H, J=7.8 Hz), 8.03 (dd, 2H, $J=7.7$, 1 Hz), 7.28 (t, 2H, $J=8.1$ Hz), 6.96 $(m, 4H), 6.85$ (t, 2H, $J=7.7$ Hz), 6.77 (d, 2H, $J=8.3$ Hz), 6.74 (d, 2H, $J=8.4$ Hz), 5.77 (m, 2H, CH=), 5.18 (d, 2H, $J=17.2$ Hz, CH₂=), 5.10 (d, 2H, $J=10.4$ Hz, CH₂=), 4.66 $(s, 4H, OCH₂CO), 3.98$ (t, 4H, $J=4.7$ Hz, $OCH₂CH₂$), 3.90 (d, 4H, $J=5.5$ Hz, OCH₂CH=), 3.78 (s, 4H, NCH₂), 3.58 (t, 4H, $J=4.7$ Hz, OCH₂CH₂). Anal. Calcd for $C_{42}H_{46}N_4O_{10}$ (766.9): C 65.78; H 6.05; N 7.31. Found: C 66.01; H 5.93; N 7.22.

3.1.10.7. Compound 33a. Yield 0.2 g (44%), colorless crystals (EtOH), mp 128 °C. LCMS; $m/z = 459$ (M+1). IR: 3396, 3333, 3071, 2873, 1685, 1659, 1601, 1531, 1483, 1457, 1334, 1293, 1253, 1226, 1163, 1132, 1091, 1046, 998, 753. ¹H NMR (CDCl₃): δ 10.54 (s, 1H, NH), 8.88 (s, 1H, NH), 8.71 (dd, 1H, $J=7.7$, 1.6 Hz), 8.43 (dd, 1H, $J=7.8$, 1.3 Hz), 8.35 (dd, 1H, $J=7.9$, 1.5 Hz), 7.49 (dt, 1H, $J=8.6$, 1.5 Hz), 7.06 (m, 7H), 6.79 (d, 1H, $J=8.0$ Hz), 6.07 (m, 1H, CH=), 5.56 (m, 1H, CH=), 5.37 (d, 1H, $J=17.3$ Hz, CH₂=), 5.27 (d, 1H, J = 10.3 Hz, CH₂=), 5.04 (dd, 1H, J = 17.2, 0.8 Hz, CH₂=), 4.94 (dd, 1H, $J=10.5$, 0.6 Hz, $CH_2=$), 4.77 (s, 2H, OCH₂CO), 4.61 (d, 2H, $J=5.6$ Hz, OCH₂CH=), 4.20 (d, 2H, $J=5.3$ Hz, OCH₂CH=). Anal. Calcd for $C_{27}H_{26}N_2O_5$ (458.5): C 70.73; H 5.72; N 6.11. Found: C 70.71; H 5.82; N 6.21.

3.1.10.8. Compound 33b. Yield 0.2 g (40%), colorless crystals (EtOH), mp 126 °C. LCMS; $m/z = 503$ (M+1). IR: 3398, 3353, 3015, 2922, 2852, 1665, 1602, 1539, 1482,

1458, 1332, 1294, 1254, 1216, 1117, 1094, 1046, 996, 930, 755. ¹H NMR (CDCl₃): δ 10.54 (s, 1H, NH), 8.91 (s, 1H, NH), 8.72 (dd, 1H, $J=8.8$, 1.2 Hz), 8.43 (d, 1H, $J=7.6$ Hz), 8.34 (d, 1H, $J=8.0$ Hz), 7.50 (t, 1H, $J=8.0$ Hz), 7.09 (m, 5H), 6.95 (d, 2H, $J=8.5$ Hz), 6.84 (d, 1H, $J=8.0$ Hz), 6.06 $(m, 1H, CH=), 5.63$ $(m, 1H, CH=), 5.36$ $(d, 1H, J=$ 17.2 Hz, CH₂=), 5.27 (d, 1H, $J=10.4$ Hz, CH₂=), 5.06 (d, 1H, $J=17.2$ Hz, CH₂=), 4.98 (d, 1H, $J=10.4$ Hz, CH₂=), 4.75 (s, 2H, OCH₂CO), 4.60 (d, 2H, $J=5.5$ Hz, OCH₂CH = $\,$, 3.82 (t, 2H, J = 4.9 Hz, OCH₂CH₂), 3.71 (d, 2H, $J=5.5$ Hz, OCH₂CH=), 3.29 (t, 2H, $J=4.9$ Hz, OCH₂CH₂). Anal. Calcd for C₂₉H₃₀N₂O₆ (502.6): C 69.31; H 6.02; N 5.57. Found: C 69.05; H 5.77; N 5.78.

3.1.11. Ring closing metathesis of 15a–e, 17b,d, 21b–e, 22, 23, 30a,b, 31a,b, 32a,b and 33a,b. General procedure. A solution of each of the substrates 15a–e, 17b,d, 21b–e, 22, 23, 30a,b, 31a,b, 32a,b and 33a,b (0.2 mmol) in DCM (25 mL) and Grubbs' catalyst $(4–12 \text{ mg}, \text{ca. } 2.5–7.5 \text{ mol\%})$ of the substrate) was heated under reflux for 2–24 h ([Tables 1–3\)](#page-165-0). The reaction mixture was then mixed with silica gel (100–200 mm, 0.5 g), stirred for 30 min, filtered and the silica was extracted twice with DCM (50 mL). After removing the solvent from the DCM extract, the remaining products were analyzed by ${}^{1}H$ NMR ([Tables 1–3\)](#page-165-0). All macrocycles were characterized by their R_f . All E and Z isomers were identified by their ${}^{1}H$ and ${}^{13}C$ NMR signals in their mixture.

3.1.11.1. Compound 1a. (E and Z): Yield 162 mg (97%), colorless crystals (EtOH), mp 243–244 °C, R_f = 0.6 [EtOAc/ pet. ether (40–60) 2:1]. LCMS; $m/z = 835$ (M + 1). IR: 3394, 3356, 3068, 2929, 1688, 1663, 1600, 1532, 1482, 1455, 1330, 1294, 1250, 1222, 1119, 1092, 1046, 1008, 955, 748. ¹H NMR (CDCl₃) (*E* isomer): δ 9.97 (s, 2H, NH), 8.78 (s, 2H, NH), 8.53 (dd, 2H, $J=8.0$, 1.3 Hz), 8.11 (dd, 2H, $J=$ 7.7, 1.5 Hz), 7.80 (dd, 2H, $J=7.9$, 1.2 Hz), 7.41 (dt, 2H, $J=$ 8.0, 1.6 Hz), 7.07 (m, 4H), 6.92 (dt, 2H, $J=7.4$, 1.3 Hz), 6.79 (m, 4H), 6.70 (d, 2H, $J=7.9$ Hz), 6.43 (d, 2H, $J=$ 7.8 Hz), 6.33 (t, 2H, $J=7.8$ Hz), 6.27 (s, 2H, CH=), 4.73 (s, 4H, OCH₂CO), 4.53 (s, 4H, $CH_2CH=$), 3.52 (s, 4H, OCH₂CH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 165.7, 162.6, 156.0, 146.5, 146.2, 133.3, 132.8, 128.6, 128, 125.2, 124.8, 123.9, 122.8, 121.9, 121.5, 121.1, 120.9, 118.3, 112.2, 111.4, 109.6, 69.6, 69.1, 66.6. ¹H NMR (CDCl₃) (Z isomer): δ 10.19 (s, 2H, NH), 8.74 (s, 2H, NH), 8.57 (d, 2H, J= 8.3 Hz), 8.18 (d, 2H, $J=7.8$ Hz), 7.98 (d, 2H, $J=7.8$ Hz), 7.41 (t, 2H, $J=8.0$ Hz), 7.08 (m, 4H), 6.98 (t, 2H, $J=$ 7.8 Hz), 6.82 (m, $4H$), 6.64 (d, $2H$, $J=8.8$ Hz), 6.52 (d, $2H$, $J=7.5$ Hz), 6.33 (t, 2H, $J=7.8$ Hz), 5.89 (t, 2H, $J=3.6$ Hz, CH=), 4.75 (s, 4H, OCH₂CO), 4.61 (d, 4H, $J=3.6$ Hz, $CH_2CH=$), 3.80 (s, 4H, OCH₂CH₂). ¹³C NMR (CDCl₃) (Z isomer): d 165.7, 165.69, 155.7, 146.7, 146.6, 133.3, 132.8, 129.6, 127.8, 125.9, 124.9, 123.6, 122.8, 121.7, 121.6, 121.1, 120.9, 119.3, 113.2, 112.1, 110.9, 70.1, 67.3, 64.9. Anal. Calcd for $C_{48}H_{42}N_4O_{10}$ (834.9): C 69.06; H 5.07; N 6.71. Found: C 69.01; H 5.03; N 6.45.

3.1.11.2. Compound 1b. (E and Z): Yield 154 mg (91%), purified using column chromatography with eluent DCM–pet. ether (40–60)–EtOAc (6/4/3), colorless crystals [DCM/pet. ether (40–60)] mp 176–177 °C, R_f = 0.6 [EtOAc/ pet. ether (40–60) 2:1]. LCMS; $m/z = 849$ (M + 1). IR: 3399,

3353, 3069, 3034, 2931, 1686, 1664, 1601, 1534, 1483, 1455, 1330, 1293, 1252, 1214, 1116, 1092, 1048, 1004, 751. ¹H NMR (CDCl₃) (*E* isomer): δ 10.14 (s, 2H, NH), 8.85 (s, 2H, NH), 8.71 (d, 2H, $J=8.0$ Hz), 8.24 (m, 4H), 7.42 (t, 2H, $J=7.7$ Hz), 7.17 (m, 4H), 7.10 (t, 2H, $J=7.8$ Hz), 6.99 (d, $2H, J=8.2$ Hz), 6.81 (d, 2H, $J=8.3$ Hz), 6.72 (m, 4H), 6.47 $(m, 2H), 6.01$ (s, $2H, CH =$), 4.72 (s, $4H, OCH₂CO$), 4.48 (s, 4H, OCH₂CH=), 3.82 (t, 4H, $J=5.5$ Hz, OCH₂), 1.78 (quint, 2H, $J=5.5$ Hz, OCH₂CH₂). ¹H NMR (CDCl₃) (Z isomer): d 10.09 (s, 2H, NH), 8.83 (s, 2H, NH), 8.77 (d, 2H, $J=9.3$ Hz), 8.29 (m, 2H), 7.44 (t, 2H, $J=8.6$ Hz), 7.25 (m, 4H), 7.17 (d, 2H, $J=8.8$ Hz), 7.05 (t, 2H, $J=8.3$ Hz), 6.81 $(d, 2H, J=4 Hz)$, 6.72 (m, 4H), 6.61 (t, 2H, $J=8.8 Hz$), 6.05 $(t, 2H, J=7.8 \text{ Hz})$, 5.55 (br, 2H, CH=), 4.65 (s, 4H, OCH₂CO), 4.52 (s, 4H, OCH₂CH=), 3.91 (t, 4H, $J=$ 6.1 Hz, OCH₂), 1.78 (quint, 2H, $J=5.5$ Hz, OCH₂CH₂). Anal. Calcd for $C_{49}H_{44}N_4O_{10}$ (848.9): C 69.33; H 5.22; N 6.60. Found: C 69.11; H 5.54; N 6.31.

3.1.11.3. Compound 1c. (*E* and *Z*): Yield 91 mg (52%), colorless crystals [DCM/pet. ether $(40-60)$], mp 237 °C, R_f =0.6 [EtOAc/pet. ether (40–60) 2:1]. LCMS; $m/z = 879$ $(M+1)$. IR: 3406, 3067, 2926, 1694, 1659, 1601, 1534, 1482, 1454, 1333, 1293, 1253, 1219, 1130, 1048, 752. ¹H NMR (CDCl₃) (*E* isomer): δ 10.33 (s, 2H, NH), 8.80 (s, 2H, NH), 8.71 (d, 2H, $J=7.2$ Hz), 8.23 (dd, 2H, $J=8.0$, 1.6 Hz), 8.19 (m, 2H), 7.29 (dt, 2H, $J=8.0$, 1.5 Hz), 7.19 (t, 2H, $J=$ 7.6 Hz), 7.07 (t, 2H, $J=7.6$ Hz), 7.01 (dt, 2H, $J=7.8$, 1.3 Hz), 6.90 (d, 2H, $J=7.8$ Hz), 6.74 (m, 6H), 6.46 (m, 2H), 6.22 (t, 2H, $J=3.8$ Hz, $=$ CH), 4.72 (s, 4H, OCH₂CO), 4.60 (d, 4H, $J=3.8$ Hz, OCH₂CH=), 3.74 (t, 4H, $J=4$ Hz, OCH₂CH₂O), 3.40 (t, 4H, $J=4$ Hz, OCH₂CH₂O). ¹³C NMR (CDCl3) (E isomer): d 166, 163.1, 156.3, 147.1, 146.6, 133.4, 132.3, 129.5, 129.4, 126.4, 124.6, 124, 124, 122, 122, 121.2, 119.5, 114, 113.1, 111.6, 70.5, 70.1, 70, 68.4. ¹ H NMR (CDCl₃) (Z isomer): δ 10.26 (s, 2H, NH), 8.88 (s, 2H, NH), 8.71 (d, 2H, $J=7.2$ Hz), 8.23 (dd, 2H, $J=8.0$, 1.6 Hz), 8.19 (m, 2H), 7.39 (m, 2H), 7.20 (m, 6H), 7.05 (m, 2H), 6.74 $(m, 6H), 6.35$ $(m, 2H), 5.67$ $(t, 2H, J=3.5$ Hz, CH $=$), 4.64 (d, 4H, $J=3.5$ Hz, OCH₂CH=), 4.61 (s, 4H, OCH₂CO), 3.74 (t, 4H, $J=4$ Hz, OCH₂CH₂O), 3.40 (t, 4H, $J=4$ Hz, OCH₂CH₂O). Anal. Calcd for $C_{50}H_{46}N_4O_{11}$ (878.9): C 68.33; H 5.28; N 6.37. Found: C 67.97; H 5.06; N 6.19.

3.1.11.4. Compound 1d. (E and Z): Yield 144 mg (78%), colorless crystals [DCM/pet. ether (40–60)], mp 201– 202 °C, R_f =0.4 [EtOAc/pet. ether (40–60) 2:1]. LCMS; $m/z = 923$ (M + 1). IR: 3392, 3342, 3068, 2936, 2875, 1686, 1659, 1600, 1532, 1481, 1456, 1330, 1293, 1254, 1220, 1163, 1134, 1118, 1092, 1048, 750. ¹H NMR (CDCl₃) (*E* isomer): d 10.35 (s, 2H, NH), 8.81 (s, 2H, NH), 8.71 (dd, 2H, $J=8.1, 1.3$ Hz), 8.38 (dd, 2H, $J=7.9, 1.4$ Hz), 8.23 (dd, 2H, $J=7.7, 1.6$ Hz), 7.00 (m, 14H), 6.86 (d, 2H, $J=8.1$ Hz), 6.68 (d, 2H, $J=7.6$ Hz), 6.43 (m, 2H, $=$ CH), 4.82 (s, 4H, OCH₂CO), 4.67 (m, 4H, OCH₂CH=), 3.84 (t, 4H, $J=$ 3.9 Hz, OCH₂CH₂), 3.45 (t, 4H, $J=3.9$ Hz, OCH₂CH₂), 3.32 (s, 4H, OCH₂CH₂O). ¹³C NMR (CDCl₃) (*E* isomer): δ 166.3, 163.2, 156.1, 147.0, 146.8, 133.6, 132.6, 129.7, 128.9, 126.7, 124.6, 123.7, 123.0, 121.7, 121.4, 121.1, 120.0, 112.2, 112.1, 112.0, 111.9, 70.3, 69.5, 69.4, 69.3, 68.3.¹H NMR (CDCl₃) (Z isomer): δ 10.40 (s, 2H, NH), 8.75 $(s, 2H, NH)$, 8.71 (dd, 2H, $J=8.1$, 1.3 Hz), 8.32 (d, 2H, $J=$ 6 Hz), 8.25 (d, 2H, $J=7.7$ Hz), 7.21 (t, 2H, $J=7.3$ Hz), 7.00

(m, 14H), 6.73 (d, 2H, $J=7$ Hz), 6.09 (t, 2H, $J=3.7$ Hz, $=$ CH), 4.82 (d, 4H, $J=3.7$ Hz, OCH₂CH $=$), 4.75 (s, 4H, OCH₂CO), 3.88 (t, 4H, $J=4$ Hz, OCH₂CH₂), 3.43 (t, 4H, $J=4$ Hz, OCH₂CH₂), 3.23 (s, 4H, OCH₂CH₂O). ¹³C NMR (CDCl₃) (Z isomer): δ 166.1, 163.2, 156.1, 147.0, 146.8, 133.2, 132.4, 129.2, 128.7, 126.8, 123.9, 123.2, 121.9, 121.8, 121.6, 121.2, 120.3, 112.6, 112.5, 112.0, 70.4, 69.1, 65.0. Anal. Calcd for $C_{52}H_{50}N_4O_{12}$ (923): C 67.67; H 5.46; N 6.07. Found: C 67.69; H 5.49; N 6.12.

3.1.11.5. Compound 1e. (E): Yield 160 mg (88%), colorless crystals (EtOH), mp 229–230 °C, R_f = 0.8 [EtOAc/ pet. ether (40–60) 2:1]. LCMS; $m/z = 911$ (M + 1). IR: 3401, 3347, 3070, 3023, 2934, 1695, 1651, 1599, 1536, 1482, 1455, 1332, 1295, 1250, 1220, 1163, 1138, 1116, 1092, 1047, 1008, 751. ¹H NMR (DMSO- d_6): δ 10.31 (s, 2H, NH), 9.50 (s, 2H, NH), 8.39 (d, 2H, $J=7.7$ Hz), 8.02 (d, 2H, $J=$ 7.4 Hz), 7.92 (d, 2H, J=7.4 Hz), 7.30 (m, 2H), 7.26 (t, 2H, $J=7.3$ Hz), 7.08 (m, 14H), 6.93 (d, 4H, $J=7.3$ Hz), 6.02 (br, 2H, $=CH$), 5.15 (s, 4H, OCH₂Ar), 4.83 (s, 4H, OCH₂CO), 4.59 (s, 4H, OCH₂CH=). ¹³C NMR (DMSO-d₆) δ = 39.5): δ 166.5, 162.4, 155.9, 148.8, 147.2, 134.4, 133.3, 131.5, 129.1, 128.1, 127.9, 127.8, 126.4, 125.4, 124.1, 122.7, 121.6, 121.5, 121.2, 120.8, 120.7, 113.3, 112.9, 112.3, 68.7, 67.9, 67.7. ¹H NMR (DMSO- d_6): δ 10.41 (s, $2H, NH$), 9.66 (s, 2H, NH), 8.50 (d, 2H, $J=7.7$ Hz), 8.02 (d, $2H, J=7.4$ Hz), 7.92 (d, $2H, J=7.4$ Hz), 7.30 (m, $2H$), 7.26 $(t, 2H, J=7.3 \text{ Hz})$, 7.08 (m, 14H), 6.93 (d, 4H, $J=7.3 \text{ Hz}$), 5.57 (br, 2H, $=$ CH), 5.15 (s, 4H, OCH₂Ar), 4.83 (s, 4H, OCH₂CO), 4.82 (br, 4H, OCH₂CH=). Anal. Calcd for $C_{54}H_{46}N_{4}O_{10}$ (910.9): C 71.20; H 5.09; N 6.15. Found: C 70.45; H 5.12; N 6.33.

3.1.11.6. Compound 2b. (E and Z): Yield 101 mg (70%), colorless crystals mp $217-218$ °C, purified using column chromatography using DCM/pet. ether (40–60)/EtOAc, $R_f = 0.4$ [EtOAc/pet. ether (40–60) 2:1]. LCMS; $m/z = 725$ $(M+1)$. IR: 3396, 3379, 3067, 2960, 2860, 1691, 1602, 1529, 1487, 1456, 1331, 1291, 1253, 1200, 1093, 1047, 750. ¹H NMR (CDCl₃) (*E* isomer): δ 8.86 (s, 2H, NH), 8.79 (s, 2H, NH), 8.51 (dd, 2H, $J=7.2$, 1.4 Hz), 8.40 (dd, 2H, $J=7$, 2 Hz), 7.07 (m, 10H), 6.66 (d, 2 H, $J=8.5$ Hz), 5.68 (s, 2 H, CH=), 4.67 (s, 4H, OCH₂CO), 3.98 (t, 4H, $J=5.6$ Hz, OCH_2CH_2), 3.78 (s, 4H, OCH_2CO), 3.65 (s, 4H, OCH₂CH=), 2.00 (quint, 2H, $J=5.6$ Hz, OCH₂CH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 167.3, 165.6, 147.0, 146.0, 128.4, 128.2, 126.4, 124.9, 124.4, 123.9, 121.5, 120.2, 120.0, 114.2, 110.8, 70.9, 70.4, 69.9, 64.1, 29.0. ¹H NMR (CDCl₃) (Z isomer): δ 8.86 (s, 2H, NH), 8.72 (s, 2H, NH), 8.40 (dd, 2H, $J=7$, 2 Hz), 8.32 (d, 2H, $J=7.38$ Hz), 7.07 (m, 10H), 6.69 (d, 2H, $J=8.1$ Hz), 5.44 (t, 2H, $J=3.8$ Hz, CH=), 4.72 (s, 4H, OCH₂CO), 3.98 (m, 8H, OCH₂CH₂, OCH₂CH=), 3.89 (s, 4H, OCH₂CO), 2.00 (quint, 2H, $J=$ 5.6 Hz, OCH₂CH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 167.1, 165.6, 147.4, 146.1, 128.3, 127.6, 126.2, 125.0, 124.6, 123.5, 121.0, 120.7, 120.5, 113.5, 110.7, 69.7, 69.6, 66.7, 64.3, 28.9. Anal. Calcd for $C_{39}H_{40}N_4O_{10}$ (724.8): C 64.63; H 5.56; N 7.73. Found: C 64.48; H 5.74; N 7.78.

3.1.11.7. Compound 2d. (E and Z): Yield 96 mg (60%), colorless crystals mp $198-199$ °C, purified using column chromatography using DCM/pet. ether (40–60)/EtOAc, $R_f = 0.1$ [EtOAc/pet. ether (40–60) 2:1]. LCMS; $m/z = 799$ $(M+1)$. IR: 3386, 2920, 1687, 1601, 1533, 1454, 1254, 1202, 1117, 1052, 955, 750. ¹H NMR (CDCl₃) (*E* isomer): δ 9.04 (s, 2H, NH), 8.77 (s, 2H, NH), 8.40 (m, 2H), 8.35 (d, 2H, $J=7.6$ Hz), 7.02 (m, 10H), 6.86 (d, 2H, $J=8.2$ Hz), 5.87 (s, 2H, CH=), 4.77 (s, 4H, OCH₂CO), 4.05 (m, 12H, OCH₂CH₂, OCH₂CH = 0CH₂CO), 3.63 (t, 4H, $J = 4.3$ Hz, OCH_2CH_2), 3.50 (s, 4H, OCH_2CH_2O). ¹H NMR (CDCl₃) (Z isomer): δ 9.05 (s, 2H, NH), 8.96 (s, 2H, NH), 8.40 (m, 2H), 8.35 (d, 2H, $J=7.6$ Hz), 7.02 (m, 10H), 6.86 (d, 2H, $J=$ 8.2 Hz), 5.70 (t, 2H, $J=3.9$ Hz, CH=), 4.76 (s, 4H, OCH₂CO), 4.19 (d, 4H, $J=3.9$ Hz, OCH₂CH=), 4.05 (m, 8H, OCH₂CH₂, OCH₂CO), 3.59 (t, 4H, $J=4.3$ Hz, OCH_2CH_2), 3.49 (s, 4H, OCH_2CH_2O). Anal. Calcd for $C_{42}H_{46}N_4O_{12}(798.9)$: C 63.15; H 5.80; N 7.01. Found: C 63.32; H 5.85; N 7.60.

3.1.11.8. Compound 3b. (*E* and *Z*): Yield 0.127 g (99%), colorless crystals (EtOH), mp $79-80$ °C, purified using column chromatography with eluent DCM/pet. ether (40– 60)/EtOAc, $R_f = 0.7$ [DCM/pet. ether (40–60)/EtOAc 6:4:2]. LCMS; $m/z = 639$ (M+1). IR: 3346, 2928, 2862, 1659, 1598, 1531, 1480, 1453, 1291, 1233, 1047, 752. ¹H NMR (CDCl₃) (*E* isomer): δ 10.11 (s, 2H, NH), 8.47 (d, 2H, $J=7.8$ Hz), 8.30 (dd, 2H, $J=7.8$, 1.4 Hz), 7.51 (dt, 2H, $J=$ 7.6, 1.6 Hz), 7.18 (t, 2H, $J=7.6$ Hz), 6.99 (m, 6H), 6.88 (d, 2H, $J=8.0$ Hz), 5.58 (s, 2H, CH=), 4.34 (m, 8H, OCH₂CH₂), 3.87 (br, 4H, OCH₂CH=), 3.77 (m, 4H, OCH₂), 2.33 (quint, 2H, $J=6$ Hz, OCH₂CH₂CH₂O). ¹³C NMR (CDCl₃) (*E* isomer): δ 163.4, 156.5, 148.2, 132.9, 132.6, 129.2, 128.3, 124.1, 123.2, 122.2, 122.0, 121.2, 113.8, 111.7, 71.1, 69.3, 67.8, 65.4, 28.8. ¹H NMR (CDCl₃) (Z isomer): δ 10.09 (s, 2H, NH), 8.51 (d, 2H, J=7.9 Hz), 8.30 (dd, 2H, $J=7.8$, 1.4 Hz), 7.49 (t, 2H, $J=7.5$ Hz), 7.18 $(t, 2H, J=7.6 \text{ Hz})$, 6.99 (m, 6H), 6.89 (d, 2H, $J=8.0 \text{ Hz}$), 5.56 (t, 2H, $J=4.2$ Hz, CH=), 4.34 (m, 4H, OCH₂), 4.27 (t, 4H, $J=4.5$ Hz, OCH₂), 4.04 (d, 4H, $J=4.2$ Hz, OCH₂CH=), 3.77 (m, 4H, OCH₂), 2.41 (quint, 2H, $J=$ 6.4 Hz, OCH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 163.4, 156.5, 148.0, 133.0, 132.5, 129.7, 128.2, 124.2, 122.9, 122.1, 121.9, 121.4, 113.6, 111.6, 69.2, 67.7, 66.3, 65.4, 29.1. Anal. Calcd for $C_{37}H_{38}N_2O_8$ (638.7): C 69.58; H 6.00; N 4.39. Found: C 69.56; H 6.13; N 4.25.

3.1.11.9. Compound 3c. (E and Z): Yield 110 mg (85%), colorless oil, purified using column chromatography with eluent DCM/pet. ether (40–60)/EtOAc, $R_f = 0.5$ [DCM/pet. ether (40–60)/EtOAc 6:4:2]. LCMS; $m/z = 669$ (M+1). IR: 3343, 2926, 2872, 1658, 1598, 1532, 1480, 1454, 1221, 1130, 1092, 1047, 771. ¹H NMR (CDCl₃) (*E* isomer): δ 10.21 (s, 2H, NH), 8.55 (m, 2H), 8.29 (m, 2H), 7.51 (dt, 2H, $J=8.2, 1.1$ Hz), 7.17 (t, 2H, $J=7.5$ Hz), 7.07 (m, 6H), 6.91 $(m, 2H), 5.70$ (br, 2H, CH=), 4.41 (t, 4H, $J=4.9$ Hz, OCH₂CH₂), 4.27 (t, 4H, $J=4.9$ Hz, OCH₂CH₂), 3.96 (br, 4H, OCH₂CH=), 3.91 (t, 4H, J=4.9 Hz, OCH₂CH₂), 3.86 (t, 4H, $J=4.9$ Hz, OCH₂CH₂). ¹³C NMR (CDCl₃) (E isomer): d 163.4, 156.6, 148.1, 132.9, 132.5, 129.3, 128.6, 124.0, 123.1, 122.0, 121.8, 121.4, 113.8, 112.2, 71.1, 69.5, 69.3, 68.2, 67.8. ¹H NMR (CDCl₃) (Z isomer): δ 10.26 (s, 2H, NH), 8.55 (m, 2H), 8.29 (m, 2H), 7.51 (dt, 2H, $J=8.2$, 1.1 Hz), 7.17 (t, 2H, $J=7.5$ Hz), 7.07 (m, 6H), 6.91 (m, 2H), 5.55 (t, 2H, $J=4$ Hz, CH=), 4.38 (t, 4H, $J=4.9$ Hz, OCH₂CH₂), 4.27 (t, 4H, $J=4.9$ Hz, OCH₂CH₂), 4.08 (d, 4H, $J=4$ Hz, OCH₂CH=), 3.91 (t, 4H, $J=4.9$ Hz,

OCH₂CH₂), 3.86 (t, 4H, $J=4.9$ Hz, OCH₂CH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 163.3, 156.7, 147.9, 133.0, 132.5, 129.4, 128.7, 123.9, 123.0, 122.0, 121.8, 121.6, 113.8, 112.4, 69.7, 69.2, 68.4, 68.0, 66.7. Anal. Calcd for $C_{38}H_{40}N_2O_9$ (668.7): C 68.25; H 6.03; N 4.19. Found: C 68.56; H 5.92; N 4.31.

3.1.11.10. Compound 3d. (E and Z): Yield 140 mg (99%), colorless oil, purified using column chromatography with eluent DCM/pet. ether $(40-60)/E$ tOAc, $R_f=0.4$ [DCM/pet. ether (40–60)/EtOAc 6:4:2]. LCMS; $m/z = 713$ (M+1). IR: 3340, 2924, 2872, 1659, 1598, 1533, 1480, 1454, 1292, 1234, 1119, 1048, 753. ¹H NMR (CDCl₃) (*E* isomer): δ 10.27 (s, 2H, NH), 8.58 (d, 2H, J=7.2 Hz), 8.28 (dd, 2H, $J=8.0$, 1.6 Hz), 7.48 (t, 2H, $J=7.8$ Hz), 7.15 (t, 2H, $J=7.6$ Hz), $7.10-7.02$ (m, 6H), 6.95 (m, 2H), 5.74 (s, 2H, CH=), 4.42 (t, 4H, $J=5.3$ Hz, OCH₂), 4.25 (t, 4H, $J=$ 5.0 Hz, OCH₂), 3.99 (br, 4H, OCH₂CH=), 3.87 (m, 8H, OCH₂), 3.67 (s, 4H, OCH₂). ¹³C NMR (CDCl₃) (*E* isomer): d 163.3, 156.6, 148.0, 132.9, 133.2, 129.2, 128.7, 123.8, 123.1, 121.9, 121.6, 121.5, 113.7, 112.1, 71.0, 70.6, 69.6, 69.0, 68.3, 67.9. ¹H NMR (CDCl₃) (Z isomer): δ 10.31 (s, 2H, NH), 8.58 (d, 2H, $J=7.2$ Hz), 8.28 (dd, 2H, $J=8.0$, 1.6 Hz), 7.48 (t, 2H, C), 7.15 (t, 2H, $J=7.6$ Hz), 7.10–7.02 $(m, 6H), 6.95$ $(m, 2H), 5.55$ $(t, 2H, J=4.0$ Hz, CH $=$), 4.41 $(t, 4H, J=5.3 \text{ Hz}, \text{ OCH}_2)$, 4.26 $(t, 4H, J=4.8 \text{ Hz}, \text{ OCH}_2)$, 4.08 (d, 4H, $J=4.0$ Hz, OCH₂CH=), 3.87 (m, 8H, OCH₂), 3.67 (s, 4H, OCH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 163.3, 156.6, 147.9, 133.0, 133.2, 129.3, 128.7, 123.8, 123.0, 121.9, 121.7, 121.6, 113.7, 112.1, 70.6, 69.7, 69.1, 68.4, 68.1, 66.8. Anal. Calcd for $C_{40}H_{44}N_2O_{10}$ (712.8): C 67.40; H 6.22; N 3.93. Found: C 67.55; H 6.49; N 3.98.

3.1.11.11. Compound 3e. (E and Z): Yield 139 mg (98%), colorless crystals (EtOH), mp 114-115 °C, purified using column chromatography with eluent DCM/pet. ether (40–60)/EtOAc, R_f =0.7 [DCM/pet. ether (40–60)/EtOAc 6:4:2]. LCMS; $m/z=701$ (M+1). IR: 3350, 2934, 2872, 1659, 1599, 1534, 1454, 1293, 1230, 1121, 1023, 751. ¹H NMR (CDCl₃) (*E* isomer): δ 10.51 (s, 2H, NH), 8.59 (d, 2H, $J=8.0$ Hz), 8.36 (d, 2H, $J=7.6$ Hz), 7.55 (m, 4H), 7.32 (m, 2H), 7.20 (t, 2H, $J=7.5$ Hz), 7.04 (d, 2H, $J=8.3$ Hz), 6.95 $(t, 2H, J=7.8 \text{ Hz})$, 6.90 (d, 2H, $J=8.3 \text{ Hz}$), 6.72 (t, 2H, $J=$ 7.8 Hz), 5.57 (s, 2H, CH=), 5.48 (s, 4H, OCH₂Ar), 4.08 (m, 4H, OCH₂), 3.82 (br, 4H, OCH₂CH=), 3.59 (m, 4H, OCH₂). ¹H NMR (CDCl₃) (Z isomer): δ 10.47 (s, 2H, NH), 8.59 (d, 2H, $J=8.0$ Hz), 8.36 (d, 2H, $J=7.6$ Hz), 7.55 (m, 4H), 7.36 (m, 2H), 7.20 (t, 2H, $J=7.5$ Hz), 7.00 (d, 2H, $J=$ 8.2 Hz), 6.95 (t, 2H, $J=7.8$ Hz), 6.90 (d, 2H, $J=8.3$ Hz), 6.84 (t, 2H, $J=7.5$ Hz), 5.48 (t, 2H, $J=4.0$ Hz, CH $=$), 5.43 (s, 4H, OCH₂Ar), 4.08 (m, 4H, OCH₂), 3.89 (d, 4H, $J=$ 4.0 Hz, $OCH_2CH = 3.59$ (m, 4H, OCH_2). Anal. Calcd for $C_{42}H_{40}N_2O_8$ (700.8): C 71.99; H 5.75; N 4.00. Found: C 71.86; H 5.65; N 3.98.

3.1.11.12. Compound 4. (E and Z): Yield 84 mg (96%), colorless crystals (EtOH), mp 203–204 °C. LCMS; $m/z =$ 441 (M+1). IR: 3367, 3073, 3018, 2919, 2850, 1650, 1601, 1533, 1482, 1450, 1298, 1241, 1117, 910, 738. ¹ H NMR (CDCl₃) (*E* isomer): δ 8.37 (br, 2H, NH), 8.25 (dd, 2H, J= 7.6, 1.3 Hz), 7.44 (m, 2H), 7.12 (t, 2H, $J=7.3$ Hz), 6.97 (d, 2H, $J=8.2$ Hz), 5.95 (br, 2H, CH=), 4.30 (m, 4H, OCH₂), 4.10 (br, 4H, OCH₂CH=), 3.88 (m, 4H, OCH₂), 3.67

(m, 4H, NCH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 165.2, 156.5, 132.5, 132.4, 128.7, 122.6, 121.9, 113.2, 71.0, 68.6, 67.9, 39.1. ¹H NMR (CDCl₃) (Z isomer): δ 8.37 (s, 2H, NH), 8.19 (dd, 2H, $J=7.8$, 1.4 Hz), 7.43 (m, 2H), 7.11 (t, 2H, $J=$ 7.3 Hz), 6.94 (d, 2H, $J=8.2$ Hz), 5.80 (t, 2H, $J=3.9$ Hz, CH=), 4.27 (m, 4H, OCH₂), 4.18 (d, 2H, $J=3.9$ Hz, OCH₂CH=), 3.88 (m, 4H, OCH₂), 3.62 (m, 4H, NCH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 165.3, 156.6, 132.4, 132.3, 128.3, 122.5, 121.7, 11.8, 68.7, 68.1, 67.0, 39.2. Anal. Calcd for $C_{24}H_{28}N_2O_6$ (440.5): C 65.44; H 6.41; N 6.36. Found: C 65.33; H 6.39; N 6.54.

3.1.11.13. Compound 5. (E and Z): Yield 57 mg (60%), colorless crystals (CH₂Cl₂), mp 208–209 °C, purified using column chromatography with eluent DCM/pet. ether $(40–60)$ /EtOAc, R_f =0.3 [DCM/pet. ether $(40–60)$ /EtOAc 6:4:2]. 2:1. LCMS; $m/z = 489$ (M+1). IR: 3303, 3072, 3007, 2931, 2863, 1664, 1600, 1536, 1475, 1454, 1303, 1233, 1162, 1124, 1099, 1047, 756. ¹H NMR (CDCl₃) (E isomer): δ 9.73 (s, 2H, NH), 8.00 (dd, 2H, $J=7.7$, 1.3 Hz), 7.81 (m, 2H), 7.43 (dt, 2H, $J=7.2$, 1.7 Hz), 7.28 (m, 2H), 7.11 (t, 2H, $J=7.7$ Hz), 6.97 (d, 2H, $J=8.3$ Hz), 5.47 (s, 2H, CH=), 4.24 (t, 4H, $J=4.2$ Hz, OCH₂), 3.92 (m, 4H, OCH₂CH=), 3.71 (t, 4H, $J=4.32$ Hz, OCH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 164.8, 156.5, 132.6, 131.4, 131.2, 129.6, 125.9, 125.1, 121.6, 113.1, 71.0, 69.2, 67.2. ¹H NMR (CDCl₃) (Z isomer): δ 9.81 (s, 2H, NH), 8.11 (dd, 2H, J= 7.9, 1.5 Hz), 7.99 (m, 2H), 7.43 (dt, 2H, $J=7.2$, 1.7 Hz), 7.28 (m, 2H), 7.11 (t, 2H, $J=7.7$ Hz), 6.97 (d, 2H, $J=$ 8.3 Hz), 5.32 (t, 2H, $J=5.3$ Hz, CH=), 4.11 (t, 4H, $J=$ 4.3 Hz, CH₂), 3.96 (d, 4H, $J=5.3$ Hz, OCH₂CH=), 3.66 (t, 4H, $J=4.3$ Hz, OCH₂). ¹³C NMR (CDCl₃) (Z isomer): δ 164.1, 156.3, 132.8, 132.2, 129.0, 125.7, 124.9, 123.9, 121.9, 113.6, 69.0, 67.7, 66.0. Anal. Calcd for $C_{28}H_{28}N_2O_6$ (488.6): C 68.84; H 5.78; N 5.73. Found: C 67.22; H 5.87; N 5.85.

3.1.11.14. Compound 6a. $(E \text{ and } Z)$: Yield 846 mg (92%), colorless crystals (EtOH), mp $203-204$ °C. LCMS; $m/z = 461$ (M + 1). IR: 3393, 3066, 1674, 1599, 1537, 1502, 1488, 1453, 1387, 1335, 1292, 1248, 1204, 1189, 1125, 1044, 961, 752. ¹H NMR (CDCl₃) (*E* isomer): 8.98 (s, 2H, NH), 8.42 (dd, 2H, $J=7.9$, 1.3 Hz), 7.12 (dt, 2H, $J=8.0$, 1.5 Hz), 7.06 (m, 6H), 6.93 (d, 2H, $J=7.4$ Hz), 6.14 (s, 2H, $CH=$), 4.71 (s, 4H, OCH₂CO), 4.57 (s, 4H, OCH₂CH=). ¹³C NMR (CDCl₃) (*E* isomer): δ : 166.5, 147.9, 147.3, 127.0, 126.9, 124.6, 123.5, 121.6, 120.6, 115.5, 111.5, 70.2, 68.1. ¹H NMR (CDCl₃) (Z isomer): δ 9.07 (s, 2H, NH), 8.42 (dd, $2H, J=7.9, 1.3 Hz$, 7.06 (m, 8H), 6.93 (d, 2H, $J=7.4 Hz$), 5.94 (t, 2H, $J=4.1$ Hz, CH=), 4.69 (s, 4H, OCH₂CO), 4.65 (d, 4H, $J=4.1$ Hz, $OCH₂CH=$). Anal. Calcd for $C_{26}H_{24}N_{2}O_{6}$ (460.5): C 67.82; H 5.25; N 6.08. Found: C 67.53; H 5.37; N 6.13.

3.1.11.15. Compound 6b. (E): Yield 80 mg (73%), colorless crystals, mp $160-161$ °C, purified using column chromatography with eluent DCM/pet. ether (40–60), R_f = 0.3 [DCM/pet. ether $(40-60)$ /EtOAc 2:1:1]. LCMS: $m/z=$ 549 (M+1). IR: 3577, 3391, 2922, 2846, 1689, 1600, 1533, 1499, 1454, 1254, 1117, 1048, 750. ¹H NMR (CDCl₃) (E isomer): δ 9.26 (s, 2H, NH), 8.42 (dd, 2H, $J=8.0$, 1.5 Hz), 7.17–7.02 (m, 10H), 6.93 (d, 2H, $J=8.0$ Hz), 5.81 (s, 2H, $CH=$), 4.76 (s, 4H, OCH₂CO), 4.25 (m, 4H, OCH₂CH₂), 3.98 (br, 4H, OCH₂CH=), 3.78 (m, 4H, OCH₂CH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 166.5, 148.5, 147.5, 129.1, 127.4, 124.4, 123.8, 121.7, 120.3, 117.6, 112.2, 71.2, 70.5, 69.0, 68.3. Anal. Calcd For $C_{30}H_{32}N_2O_8$ (548.6): C 65.68; H 5.88; N 5.11. Found: C 65.92; H 6.02; N 4.99.

3.1.11.16. Compound 7a. $(E \text{ and } Z)$: Yield 116 mg (91%), colorless crystals (EtOH/CHCl₃), mp 271–272 °C. LCMS; $m/z = 637$ (M+1). IR: 3392, 3059, 2919, 1687, 1599, 1534, 1456, 1253, 1213, 1054, 1044, 966, 749. ¹H NMR (CDCl₃) (*E* isomer): δ 8.37 (s, 2H, NH), 7.94 (d, 2H, $J=8.9$ Hz), 7.90 (dd, 2H, $J=8.0$, 1.6 Hz), 7.74 (m, 2H), 7.48 (d, 2H, $J=8.9$ Hz), 7.29 (m, 4H), 7.19 (m, 2H), 7.05 $(dt, 2H, J=7.9, 1.6 Hz), 6.94 (dt, 2H, J=7.8, 1.1 Hz), 6.82$ (dd, 2H, $J=8.2$, 1.2 Hz), 5.98 (t, 2H, $J=2.2$ Hz, CH=), 4.61 (d, 2H, $J=15.4$ Hz, OCH₂CO), 4.43 (d, 4H, $J=2.2$ Hz, $CH_2CH=$), 4.37 (d, 2H, $J=15.4$ Hz, OCH₂CO). ¹³C NMR (CDCl₃) (*E* isomer): δ 166.3, 153.9, 147.6, 133.6, 130.5, 130.4, 128.0, 127.2, 127.1, 126.2, 125.1, 125.0, 124.6, 121.6, 121.2 (2C), 117.7, 111.3, 71.6, 67.9. ¹ H NMR (CDCl₃) (Z isomer): δ 8.70 (s, 2H, NH), 8.09 (dd, 2H, J= 7.9, 1.8 Hz), 8.00 (d, 2H, $J=8.8$ Hz), 7.74 (m, 2H), 7.49 (d, $2H, J=8.8$ Hz), 7.41 (t, $2H, J=7.6$ Hz), 7.29 (m, $2H$), 7.19 $(m, 2H), 7.13$ (d, 2H, $J=8.6$ Hz), 6.98 (dt, 2H, $J=7.8$, 1.5 Hz), 6.82 (dd, 2H, $J=8.2$, 1.2 Hz), 5.47 (t, 2H, $J=4$ Hz, $CH=$), 4.52 (d, 2H, $J=15.9$ Hz, OCH₂CO), 4.49 (d, 4H, $J=4$ Hz, $CH_2CH=$), 4.40 (d, 2H, $J=15.9$ Hz, OCH₂CO). ¹³C NMR (CDCl₃) (Z isomer): δ 166.8, 153.2, 147.8, 133.9, 130.4, 130.1, 128.4, 128.2, 127.1, 127.0, 125.5, 124.8, 124.7, 121.9, 121.5, 120.1, 115.8, 112.9, 69.8, 64.9. Anal. Calcd for $C_{40}H_{32}N_2O_6$ (636.7): C 75.46; H 5.07; N 4.40. Found: C 75.64; H 5.02; N 4.45.

3.1.11.17. Compound 7b. $(E \text{ and } Z)$: Yield 140 mg (97%), colorless crystals (EtOH), mp $212-213$ °C. LCMS; $m/z = 725$ (M + 1). IR: 3397, 2922, 1707, 1686, 1535, 1456, 1362, 1255, 1219, 1117, 912, 753. ¹H NMR (CDCl₃) (E isomer): δ 8.73 (s, 2H, NH), 8.17 (dd, 2H, $J=7.8$, 1.6 Hz), 8.00 (d, 2H, $J=8.9$ Hz), 7.88 (d, 2H, $J=8.2$ Hz), 7.47 (d, $2H, J=8.9$ Hz), 7.37 (dt, $2H, J=8.1$, 1.1 Hz), 7.27 (dt, $2H$, $J=7.1$, 1.2 Hz), 7.16 (d, 2H, $J=8.4$ Hz), 7.03 (dt, 2H, $J=$ 7.8, 1.6 Hz), 6.96 (dt, 2H, $J=7.8$, 1.2 Hz), 6.84 (dd, 2H, $J=8.1, 1.3$ Hz), 5.33 (t, 2H, $J=2.8$ Hz, CH $=$), 4.65 (d, 2H, $J=15.6$ Hz, OCH₂CO), 4.33 (d, 2H, $J=15.6$ Hz, OCH₂CO), 4.00 (m, 4H, OCH₂CH₂), 3.60 (dd, 2H, $J=$ 12.2, 2.8 Hz, OCH₂CH= $)$, 3.51 (m, 2H, OCH₂CH₂), 3.40 (dd, 2H, $J=12.2$, 2.8 Hz, OCH₂CH=), 3.24 (m, 2H, OCH₂CH₂). ¹³C NMR (CDCl₃) (*E* isomer): δ 166.9, 153.8, 147.8, 133.8, 130.5, 130.3, 128.9, 128.1, 127.3, 127.1, 125.5, 124.7, 124.3, 121.6, 120.9, 120.5, 117.2, 112.6, 68.3, 69.2, 71.0, 71.1. ¹H NMR (CDCl₃) (Z isomer): δ 8.64 (s, 2H, NH), 8.13 (dd, 2H, $J=8.0$, 1.6 Hz), 8.01 (d, 2H, $J=8.9$ Hz), 7.85 (d, 2H, $J=8.2$ Hz), 7.49 (d, 2H, $J=$ 8.9 Hz), 7.38 (dt, 2H, $J=8.1$, 1.1 Hz), 7.27 (dt, 2H, $J=6.8$, 1.2 Hz), 7.16 (d, 2H, $J=8.4$ Hz), 7.03 (dt, 2H, $J=7.8$, 1.6 Hz), 6.96 (dt, 2H, $J=7.8$, 1.2 Hz), 6.82 (dd, 2H, $J=8.1$, 1.3 Hz), 5.43 (t, 2H, $J=3.5$ Hz, CH=), 4.63 (d, 2H, $J=$ 15.6 Hz, OCH₂CO), 4.45 (d, 2H, $J=15.6$ Hz, OCH₂CO), 4.00 (m, 4H, OCH₂CH₂), 3.67 (dd, 2H, $J=12.2$, 3.5 Hz, OCH₂CH=), 3.53 (dd, 2H, $J=12.2$, 3.5 Hz, OCH₂CH=), 3.51 (m, 2H, OCH₂CH₂), 3.24 (m, 2H, OCH₂CH₂).¹³C NMR (CDCl₃) (Z isomer): δ 166.6, 153.8, 147.8, 133.7, 130.3, 130.0, 128.9, 128.1, 127.1, 126.8, 125.3, 124.8,

124.6, 121.4, 121.3, 120.6, 116.9, 112.0, 70.9, 69.0, 68.0, 66.9. Anal. Calcd for $C_{44}H_{40}N_2O_8$ (724.8): C 72.91; H, 5.56; N, 3.86. Found: C 71.97; H 5.49; N 4.05.

3.1.11.18. Compound 8a. (E and Z): Yield 130 mg (100%), colorless crystals (EtOH), mp $205 \degree C$. LCMS; $m/z = 651$ (M + 1). IR: 3396, 3351, 3070, 2938, 2867, 1691, 1649, 1601, 1535, 1484, 1454, 1374, 1292, 1249, 1206, 1161, 1116, 1043, 754. ¹H NMR (CDCl₃) (*E* isomer): δ 9.10 $(s, 2H, NH)$, 8.65 $(s, 2H, NH)$, 8.36 (dd, $2H, J=8.2, 1.7 Hz$), 8.03 (d, 2H, $J=7.6$ Hz), 7.30 (m, 2H), 7.03 (m, 4H), 6.93 (d, 2H, $J=8.3$ Hz), 6.88–6.75 (m, 4H), 5.60 (s, 2H, CH=), 4.94 (s, 4H, OCH₂CO), 4.21 (s, 4H, OCH₂CH=), 3.72 (s, 4H, CH₂N). ¹³C NMR (CDCl₃) (*E* isomer): δ 165.9, 165.6, 155.2, 147.5, 133.1, 132.4, 127.3, 124.7, 122.2, 121.5, 120.4, 112.1, 111.9, 68.5, 68.3, 41.4. ¹H NMR (CDCl₃) (Z isomer): δ 8.89 (s, 2H, NH), 8.73 (s, 2H, NH), 8.44 (m, 2H), 8.08 (d, 2H, $J=8.3$ Hz), 7.33 (m, 2H), 7.03 (m, 4H), 6.89– 6.77 (m, 2H), 6.66 (m, 2H), 5.75 (t, 2H, $J=3.4$ Hz, CH=), 4.94 (s, 4H, OCH₂CO), 4.24 (d, 4H, $J=3.4$ Hz, OCH₂CH = \vert , 3.84 (s, 4H, CH₂N). ¹³C NMR (CDCl₃) (Z isomer): d 165.9, 165.6, 155.2, 146.8, 133.1, 132.4, 127.3, 124.7, 122.2, 121.5, 120.4, 112.1, 111.9, 68.5, 68.3, 41.4. Anal. Calcd for $C_{36}H_{34}N_4O_8$ (650.7): C 66.45; H 5.27; N 8.61. Found: C 66.19; H 5.45; N 8.34.

3.1.11.19. Compound 8b. $(E \text{ and } Z)$: Yield 118 mg (80%), colorless crystals [EtOAc/pet. ether (40–60)], mp 174–175 °C. LCMS; $m/z = 739$ (M+1). IR: 3397, 3353, 2931, 2872, 1687, 1642, 1602, 1539, 1484, 1454, 1299, 1255, 1208, 1162, 1118, 1045, 909, 732, 649. ¹ H NMR (CDCl₃) (*E* isomer): δ 8.71 (s, 2H, NH), 8.60 (s, 2H, NH), 8.28 (dd, 2H, $J=7.7$, 1.2 Hz), 8.04 (d, 2H, $J=7.7$ Hz), 7.28 $(t, 2H, J=7.7 \text{ Hz})$, 7.10–6.80 (m, 10H), 5.74 (s, 2H, CH=), 4.83 (s, 4H, OCH2CO), 4.13 (m, 4H, OCH2), 3.89 (s, 4H, OCH₂CH=), 3.82 (m, 4H, OCH₂CH₂), 3.50 (s, 4H, OCH₂N). ¹³C NMR (CDCl₃) (*E* isomer): δ 166.2, 164.9, 155.5, 147.3, 132.8, 132.0, 128.7 (2C), 127.5, 124.5, 122.3, 122.0, 120.8, 113.1, 112.5, 70.9, 68.7, 68.6, 68.5, 40.5. ¹H NMR (CDCl₃) (*Z* isomer): δ 8.83 (s, 2H, NH), 8.51 (s, 2H, NH), 8.28 (dd, 2H, $J=7.7$, 1.2 Hz), 8.04 (d, 2H, $J=7.7$ Hz), 7.35 (t, 2H, $J=8.4$ Hz), 7.10–6.80 (m, 10H), 5.52 (t, 2H, $J=4.2$ Hz, CH=), 4.85 (s, 4H, OCH₂CO), 4.13 (m, 8H, OCH_2CH_2 , $OCH_2CH = 0.3.82$ (m, 4H, OCH_2CH_2), 3.50 (m, 4H, OCH₂N). ¹³C NMR (CDCl₃) (Z isomer): δ 166.1, 166.0, 155.7, 147.7, 132.9, 132.1, 129.1, 128.7 (2C), 124.8, 122.5, 122.3, 120.9, 113.1, 112.5, 69.3, 69.0, 68.4, 66.7, 40.8. Anal. Calcd for $C_{40}H_{42}N_4O_{10}$ (738.8): C 65.03; H 5.73; N 7.58. Found: C 65.45; H 5.54; N 8.37.

3.1.11.20. Compound 9a. (E and Z): Yield 65 mg (75%), colorless crystals [EtOAc/pet. ether $(40-60)$], mp 192 °C. LCMS; $m/z = 431 (M+1)$. IR: 3398, 1690, 1662, 1599, 1531, 1483, 1455, 1329, 1294, 1254, 1235, 1043, 1002, 966, 750. ¹H NMR (CDCl₃) (*E*): δ 9.80 (s, 1H, NH), 8.88 (s, 1H, NH), 8.49 (m, 1H), 8.29 (m, 2H), 7.53 (dt, 1H, $J=7.8$, 1.2 Hz), 7.18 (t, 1H, $J=7.6$ Hz), 7.10–7.04 (m, 5H), 6.96 (m, 1H), 6.90 (dd, 1H, $J=7.8$, 1.0 Hz), 6.47 (m, 1H, CH=), 6.17 (m, 1H, CH=), 4.77 (br, 2H, OCH₂CH=), 4.76 (s, 2H, OCH₂CO), 4.60 (br, 2H, OCH₂CH=). ¹H NMR (CDCl₃) (Z) : δ 9.80 (s, 1H, NH), 9.00 (s, 1H, NH), 8.40 (m, 2H), 8.23 $(m, 1H), 7.50$ (t, $1H, J=7.8$ Hz), $7.18-7.04$ (m, 6H), 6.96 (m, 2H), 6.45 (t, 2H, $J=5.3$ Hz, CH=), 4.89 (d, 2H, $J=5.3$ Hz,

 $OCH_2CH=$), 4.71 (s, 2H, OCH₂CO), 4.62 (d, 2H, $J=5.3$ Hz, OCH₂CH=). Anal. Calcd for C₂₅H₂₂N₂O₅ (430.5): C 69.76; H 5.15; N 6.51. Found: C 60.86; H 5.18; N 6.55.

3.1.11.21. Compound 9b. (E and Z): Yield 78 mg (82%), colorless crystals (EtOH/CHCl₃), mp 215–216 °C. LCMS; $m/z = 475$ (M + 1). IR: 3342, 2919, 1667, 1599, 1533, 1482, 1455, 1294, 1229, 1120, 1092, 1007, 911, 746. ¹ H NMR $(CDCl₃)$: (E isomer): 10.55 (s, 1H, NH), 9.04 (s, 1H, NH), 8.63 (dd, 1H, $J=8.6$, 0.9 Hz), 8.35 (m, 2H), 7.52 (t, 1H, $J=$ 7.4 Hz), 7.19–6.96 (m, 8H), 6.21 (m, 1H, CH=), 5.97 (m, 1H, CH=), 4.78 (s, 2H, OCH₂CH=), 4.66 (s, 2H, OCH₂CO), 4.24 (br, 2H, OCH₂CH₂), 4.02 (s, 2H, $OCH_2CH =$), 3.67 (br, 2H, OCH_2CH_2). ¹H NMR (CDCl₃) (Z isomer): d 10.55 (s, 1H, NH), 8.82 (s, 1H, NH), 8.73 (d, $1H, J=7.8$ Hz), 8.38 (m, 2H), 7.52 (t, 1H, $J=7.3$ Hz), 7.19– 6.96 (m, 7H), 6.87 (d, 1H, $J=7.7$ Hz), 5.97 (m, 2H, CH=), 4.87 (d, 2H, $J=5.6$ Hz, OCH₂CH=), 4.80 (s, 2H, OCH₂CO), 4.26 (d, 2H, $J=5.6$ Hz, OCH₂CH=), 4.08 (br, $2H$, OCH₂CH₂), 3.66 (br, 2H, OCH₂CH₂). Anal. Calcd for $C_{27}H_{26}N_2O_6$ (474.5): C 68.34; H 5.52; N 5.90. Found: C 67.41; H 5.40; N 6.06.

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A one-pot access to cycloalkano[1,2-a]indoles through an intramolecular alkyl migration reaction in indolylborates

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Abstract—A novel one-pot protocol for the preparation of cycloalkano[1,2-a]indoles by way of an intramolecular alkyl migration reaction in cyclic indolylborates is described. NaOMe was found to act as a successful trialkylboryl-protecting group against to the lithiation at the C2 of the indole ring. Treatment of cyclic indolylborates with electrophiles produced cycloalkano-[1,2-a]indoles. $© 2005 Elsevier Ltd. All rights reserved.$

1. Introduction

The rich chemistry of organoboron compounds has provided fertile ground for the development of pivotal synthetic methodologies.^{[1](#page-185-0)} While an inter- or intramolecular alkyl migration from boron to carbon in organoboron compounds has been well recognized as a valuable synthetic tool for regioselective and stereospecific bond formation, 2 its use for intramolecular cyclization has been scarcely known.^{[3](#page-185-0)} In our continuing program to develop trialkyl $(1H$ -indol-2-yl)borate as a versatile synthetic intermediate for the construction of indole derivatives,^{[4](#page-185-0)} an intramolecular alkyl migration from boron to the C2 of the indole ring in indolylborates has also been proven to be successful, leading to 2,3-disubstituted indoles in a one-pot treatment. Hence, we have become interested in the unprecedented use of the alkyl migration process in indolylborates, and previously reported a novel one-pot protocol for the preparation of carbazole derivatives based on the intramolecular 1,2-alkyl migration reaction in indolylborates, in which π -allyl palladium complexes were adopted as successful intra-molecular electrophiles.^{[5](#page-185-0)}

As the core structure of [a]-annelated indole is present in a number of biologically active indole derivatives such as mitomycin and vincamine, the development of methods for the construction of [a]-annelated indole nuclei has been the subject of a number of reports.^{[6](#page-185-0)} We have set about the

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development of a novel one-pot access to [a]-annelated indoles by the use of the 1,2-alkyl migration process in indolylborate (6) .^{[7](#page-186-0)}

2. Results and discussion

As shown in [Scheme 1](#page-178-0), we initially envisioned that alkyl migration triggered by intermolecular attack of electrophile $(H₂O)$ on the C-3 of the indole ring in cyclic indolylborate (6) might provide the cyclization product (8) after oxidation of 7, in which implementation of the synthetic plan first required an adequate protocol for the in situ generation of 6. Our initial expectation to form 6 via 2-lithioindole (A) involved straightforward lithiation of the starting indole (1), followed by treatment with dialkylboranes, but all attempted experiments have met with failure. Alternatively, we anticipated that if 2-lithioindole (5) is available by the lithiation at the C-2 of the indole ring of alkylborane (2), the following spontaneous cyclization might possibly provide cyclic indolylborate (6).

Initially, we attempted the lithiation with tert-BuLi at the C-2 of the indole ring in alkylboranes (2), readily generated by treatment of indole (1a) with 9-borabicyclo[3.3.1]nonane (9-BBN) in THF at room temperature. However, an oxidative work-up of the reaction mixture allowed only the isolation of alcohol (10a) in 40% yield, which possibly involved the formation of tetraalkylborate (B) from the predominant interaction between the trialkylboryl group of 2 and tert-BuLi. With this in mind, we needed a feasible trialkylboryl-protecting group that would persist until the lithiation at the C2 of 2 was complete, and after that, would

Keywords: Cyclic indolylborate; Hydroboration; Intramolecular alkyl migration; [a]-Annelated indoles.

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Scheme 1.

be removed to restore the trialkylboryl group. We were pleased to find that the presence of NaOMe provided marked protection against the lithiation of 2 by the formation of methoxyborate (3), and the following workup of the reaction mixture with H_2O_2 in NaOH solution afforded 8, accompanied by a small amount of 10. The reaction outcome can be interpreted as follows: (1) hydroboration of 1 with 9-BBN produces alkylborane (2). (2) Treatment of 2 with NaOMe forms methoxyborate (3). (3) Subsequent lithiation of 3 with tert-BuLi generates 2-lithioindole (4), which is accompanied by spontaneous formation of indolylborate (6) via elimination of the methoxide anion (4 to 5). (4) Intramolecular 1,2-alkyl migration in 6 provides $[a]$ -annelated indole (8) after oxidation of 7. Alcohols (10) were possibly produced by way of oxidation of 5 or B.

As summarized in [Table 1](#page-179-0), the use of NaOMe (1.1 equiv), tert-BuLi (2.2 equiv) and TMEDA (2.2 equiv) in THF was adequate to effect the one-pot transformation of 1a to 8a, and the conditions were applied to further investigations. Hydroboration of 5d brought about 6 $(n=1, X=5-NO₂)$ in situ, which was subsequently subjected to the reaction. However, the desired cyclization product was not obtained in this transformation and only alcohol (10b) was isolated from the complex reaction mixtures.

	NaOMe (equiv)	tert-BuLi (equiv)	TMEDA (equiv)	Yield $(\%)$ of 8	Yield $(\%)$ of 10	
$n=1$ X = H		2.2	2.2	8a(16)	10a (20)	
$n=1$ X = H	1.1	2.2	2.2	8a(62)		
$n=1$ X = H	2.2	2.2	2.2	8a(22)	10a (10)	
$n=1$ X = H	1.1	1.1	2.2	8a(14)	10a (22)	
$n=1$ X = H	1.1	3.0	2.2	8a(47)	10a (7)	
$n=1$ X = H	1.1	2.2		8a(40)	10a (10)	
$n=1$ X = H	1.1	2.2	1.1	8a(50)	10a (5)	
$n=1$ X = 7-Me	1.1	2.2	2.2	8b(53)		
$n=1$ X = 5-OMe	1.1	2.2	2.2	8c(60)		
$n=1$ X = 5-NO ₂	1.1	2.2	2.2		10 $b(40)$	
$n=1$ X = H	1.1	2.2	2.2	8d(60)	10 $c(5)$	
$n=1$ X = H	1.1	2.2	2.2	8e (42)	10 $d(8)$	

Table 1. One-pot preparation of 8 from 1^a

^a Yield $(\%)$ of 8 and 10 based on 1.

Treatment of 6 $(X=H)$ with various electrophiles such as alkyl halides and π -allyl palladium complexes similarly produced $[a]$ -annelated indoles (9) , which allowed the introduction of various functional groups at the C3 of the indole ring (Table 2). On the reaction of indolylborate (6; $n=2$, X=H) with 3-bromocyclohexene, borinate (9k) was isolated as stable crystals in 35% yield after oxidation of the reaction mixture, which was in contrast to the formation of 9*j* and 9l from the reaction of 6 ($n=1,3$, $X = H$). Longer oxidation time and use of increased amount of H_2O_2 did not effect the production of **9k**. As there are examples of the successful use of diphenyliodonium ion as an electrophile toward the enolate anion, 8 the diphenyliodonium ion was also expected to be suitable for the promotion of the alkyl migration in 6. The reaction of 6 $(X=H, n=1)$ with diphenyliodonium chloride in THF under the same conditions afforded furanylindole (9t) in 30% yield as the only isolable product. This is probably due to the alkyl migration in 6 ($n=1$, $X=H$) caused by the electrophilic attack of the furanium ion (C) arising from the rapid oxidation of THF by diphenyliodonium ion^{[9](#page-186-0)} ([Scheme 2](#page-180-0)).

The alkyl migration reaction in indolylborates (12), derived from indoles (11) having a substituent at the olefinic carbon, was next examined. Indole (11a) was successfully

Table 2. One-pot preparation of 9 from 1 $(X=H)^a$

^a Yield (%) of **9** and **10** based on **1**.
^b Compound **9k** was isolated.

Scheme 2.

transformed to 13a via 12 ($R = Me$). Otherwise, hydroboration of 11b became much more sluggish, requiring forced reaction conditions (reflux for 2 h), and the following steps were less effective in giving 13b in low yield, accompanied by a substantial amount of alcohol (14b). When indole (11c) bearing an alkoxy group was subjected to the alkyl migration reaction, 1-allyl-2-oxyindole (17) was isolated without the expected cyclization product. The rapid elimination of the alkoxy group from borate (15) predominantly took place to generate 16, and the subsequent oxidation gave rise to 17, as shown in Scheme 3.

Aqueous treatment of 19, derived from indoles (18) bearing a substituent at the C-3 of the indole ring, affordrd $[a]$ -annelated indoles (9a, 20) and alcohols (10e, 21), respectively ([Scheme 4\)](#page-181-0). In the case of 18b, LDA was employed for the

Scheme 4.

lithiation at the C-2 of the indole ring in order to generate indolylborate (19b). Otherwise, simply treating 19a with allyl bromide and the following work-up of the reaction mixture without a conventional oxidation allowed the isolation of alkylborane (22) in 35% yield as stable crystals. The reaction proceeded through the electrophilic attack of allyl bromide at the C-3 of the indole ring in 19a with simultaneous alkyl migration in an *anti* manner^{5} to produce 22, whose structure was confirmed based on NOE experiments.

In summary, we have demonstrated a new one-pot access to cycloalkano $[1,2-a]$ indoles (9) by way of the unprecedented use of an intramolecular alkyl migration process in indolylborate (6). Further extension of the protocol for the preparation of indole alkaloid is under way.

3. Experimental

3.1. General

Melting points were recorded on a Yamato MP21 and are uncorrected. MS and high-resolution MS spectra were recorded on a Micromass AutoSpec 3100 mass spectrometer. IR spectra were measured on a Hitachi Model 270-30 spectrometer. The NMR experiments were performed with a JEOL JNM-LA300 or JNM-ECA500 spectrometer, and chemical shifts are expressed in ppm (δ) with TMS as an internal reference. Medium pressure liquid chromatography (MPLC) was performed on silica gel (Silica Gel 60N, Kanto Chemical Co., Ltd).

3.2. General procedure for the preparation of 8

To a solution of 1 (2 mmol) in THF (10 mL), 9-BBN (0.5 M solution in THF, 2.2 mmol) was added at room temperature under an argon atmosphere, and the mixture was stirred for 1.5 h. Sodium methoxide (2.2 mmol) was added to the mixture, and after stirring for 30 min, TMEDA (4.4 mmol) and tert-BuLi (1.5 M solution in pentane, 4.4 mmol) were added to the mixture at -20 °C. After stirring for 2 h, the mixture was gradually raised to room temperature, and stirred for 4 h. To the reaction mixture, 20% NaOH (10 mL) and 30% $H₂O₂$ (2 mL) were added under ice-cooling and the whole was stirred for 30 min. The mixture was diluted with ethyl acetate (100 mL), washed with brine, and dried over MgSO₄. The solvent was removed, and the residue was separated by MPLC to give 8 (with hexane/AcOEt=100:1) and 10 (with hexane/ $AcOEt = 1:1$).

3.2.1. 2,3-Dihydro-1H-pyrrolo[1,2-a]indole $(8a)$. Mp 77–78 °C (lit.^{[10](#page-186-0)} 78–79 °C). ¹H NMR (CDCl₃) δ : 2.59 (tt, 2H, $J=6.8$, 7.3 Hz), 3.00 (t, 2H, $J=7.3$ Hz), 4.04 (t, 2H, $J=6.8$ Hz), 6.15 (s, 1H), 7.03 (dt, 1H, $J=1.4$, 7.1 Hz), 7.10 (dt, 1H, $J=1.4$, 7.1 Hz), 7.22 (d, 1H, $J=7.1$ Hz), 7.53 (d, 1H, $J=7.1$ Hz). ¹³C NMR (CDCl₃) δ : 24.1, 27.7, 43.4, 92.1, 109.3, 119.9, 120.1, 132.5, 133.2, 144.5. MS m/z : 157 (M⁺).

 $3.2.2.$ 5-Methyl-2.3-dihydro-1H-pyrrolo[1,2-a]indole **(8b).** Mp 90–91 °C (hexane). ¹H NMR (CDCl₃) δ : 2.64 (s, $3H$), $2.55-2.65$ (m, $2H$), 2.94 (t, $2H$, $J=7.4$ Hz), 4.36 (t, $2H$, $J=6.8$ Hz), 6.11 (s, 1H), 6.81 (d, 1H, $J=7.9$ Hz), 6.91 (t, 1H, $J=7.9$ Hz), 7.34 (d, 1H, $J=7.9$ Hz). ¹³C NMR (CDCl₃) d: 17.9, 23.7, 27.9, 46.5, 92.6, 118.1, 119.3, 120.2, 121.7, 132.3, 133.4, 144.7. MS m/z : 171 (M⁺). Anal. Calcd for $C_{12}H_{13}N$: C, 84.17; H, 7.65; N, 8.18. Found: C, 84.38; H, 7.55; N, 8.24.

3.2.3. 7-Methoxy-2,3-dihydro-1H-pyrrolo[1,2-a]indole **(8c).** Mp 85–87 °C (lit.^{[11](#page-186-0)} 84–85 °C). ¹H NMR (CDCl₃) δ : $2.50-2.60$ (m, 2H), 2.97 (t, 2H, $J=7.8$ Hz), 3.83 (s, 3H),

3.98 (t, 2H, $J=7.0$ Hz), 6.08 (s, 1H), 6.77 (d, 1H, $J=$ 7.8 Hz), 7.02 (br s, 1H), 7.09 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl3) d: 24.4, 27.7, 43.7, 55.9, 92.0, 102.6, 109.9, 128.1, 133.6, 145.3, 153.9. MS m/z : 187 (M⁺).

3.2.4. 6,7,8,9-Tetrahydropyrido[1,2-a]indole (8d). Mp 58–59 °C (lit.^{[12](#page-186-0)} 57–58 °C). ¹H NMR (CDCl₃) δ : 1.80–1.95 $(m, 2H), 2.00-2.15$ $(m, 2H), 2.97$ $(t, 2H, J=6.3$ Hz $), 3.00$ $(t,$ $2H, J=7.3$ Hz), 4.04 (t, 2H, $J=6.4$ Hz), 6.14 (s, 1H), 7.06 (dt, 1H, $J=1.5$, 7.8 Hz), 7.13 (dt, 1H, $J=1.5$, 7.8 Hz), 7.26 (d, 1H, $J=7.8$ Hz), 7.52 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl3) d: 21.1, 23.3, 24.1, 42.1, 97.3, 108.4, 119.4, 119.9, 128.1, 136.1, 137.0. MS m/z : 171 (M⁺).

3.2.5. 7,8,9,10-Tetrahydro-6H-azepino[1,2-a]indole (8e). Mp 86–85 °C (lit.^{[12](#page-186-0)} 83–85 °C). ¹H NMR (CDCl₃) δ : 2.59 (tt, 2H, $J=6.8$, 7.3 Hz), 3.00 (t, 2H, $J=7.3$ Hz), 4.04 (t, 2H, $J=6.8$ Hz), 6.15 (s, 1H), 7.03 (dt, 1H, $J=1.4$, 7.1 Hz), 7.10 (dt, 1H, $J=1.4$, 7.1 Hz), 7.22 (d, 1H, $J=7.1$ Hz), 7.53 (d, 1H, J = 7.1 Hz). ¹³C NMR (CDCl₃) δ : 8.1, 28.6, 29.4, 31.0, 44.5, 98.9, 108.5, 118.8, 119.7, 120.2, 127.7, 136.8, 143.2. MS m/z : 185 (M⁺).

3.3. General procedure for the preparation of 9

To a solution of 1 (2 mmol) in THF (10 mL), 9-BBN (0.5 M solution in THF, 2.2 mmol) was added at room temperature under an argon atmosphere, and the mixture was stirred for 1.5 h. Sodium methoxide (2.2 mmol) was added to the mixture, and after stirring for 30 min, TMEDA (4.4 mmol) and tert-BuLi (1.5 M solution in pentane, 4.4 mmol) were added to the mixture at -20 °C. After stirring for 2 h, the mixture was gradually raised to room temperature, and stirred for 4 h. Then, electrophile (5 mmol) was added, and the whole was stirred overnight (in the cases of alkyl halides) or heated under reflux for 3 h (in the cases of π -allyl palladium complexes). To the reaction mixture, 20% NaOH (10 mL) and 30% H_2O_2 (2 mL) were added under icecooling and the whole was stirred for 30 min. The mixture was diluted with ethyl acetate (100 mL), washed with brine, and dried over MgSO4. The solvent was removed, and the residue was separated by MPLC to give 9 (with hexane/ $AcOEt = 100:1-10:1$ and 10 (with hexane/AcOEt = 1:1).

3.3.1. 9-Methyl-2,3-dihydro-1H-pyrrolo[1,2-a]indole **(9a).** Mp 50–51 °C (lit.^{[13](#page-186-0)} 48–49 °C). ¹H NMR (CDCl₃) δ : 2.24 (s, 3H), 2.52 (m, 2H), 2.88 (t, 2H, $J=7.3$ Hz), 3.95 (t, $2H, J=6.9$ Hz), $7.00-7.12$ (m, $2H$), 7.15 (d, $1H, J=7.8$ Hz), 7.45 (d, 1H, $J=7.3$ Hz). ¹³C NMR (CDCl₃) δ : 8.8, 22.7, 27.5, 43.3, 100.4, 108.7, 118.1, 119.8, 133.0, 132.4, 141.1. MS m/z : 171 (M⁺).

3.3.2. 5-Methyl-6,7,8,9-tetrahydropyrido[1,2-a]indole (9b). ¹H NMR (CDCl₃) δ : 1.88–1.94 (m, 2H), 2.04–2.10 $(m, 2H), 2.24$ (s, 3H), 2.90 (t, 2H, $J=6.3$ Hz), 4.03 (t, 2H, $J=6.3$ Hz), 7.11 (t, 1H, $J=7.8$ Hz), 7.17 (t, 1H, $J=8.0$ Hz), 7.25 (d, 1H, $J=8.0$ Hz), 7.52 (d, 1H, $J=8.1$ Hz). ¹³C NMR (CDCl3) d: 8.2, 21.4, 22.5, 23.7, 42.4, 104.8, 108.4, 117.8, 119.0, 120.2, 128.6, 133.0, 136.0. HR-MS m/z: Calcd for C₁₃H₁₅N: 185.1204. Found: 185.1192.

3.3.3. 11-Methyl-7,8,9,10-tetrahydro-6H-azepino[1,2-a] indole (9c). Mp $88-89$ °C (hexane). ¹H NMR (CDCl₃)

 δ : 1.67–1.78 (m, 4H), 1.81–1.88 (m, 2H), 2.25 (s, 3H), 2.86 (t, 2H, $J=5.8$ Hz), 4.13 (t, 2H, $J=5.2$ Hz), 7.04 (t, 1H, $J=7.8$ Hz), 7.13 (t, 1H, $J=7.8$ Hz), 7.23 (d, 1H, $J=$ 8.0 Hz), 7.48 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 8.8, 25.5, 28.0, 29.8, 31.3, 44.6, 105.5, 108.2, 118.3, 120.4, 128.3, 135.8, 139.1. MS m/z : 199 (M⁺). Anal. Calcd for $C_{14}H_{17}N$: C, 84.37; H, 8.60; N, 7.03. Found: C, 84.42; H, 8.77; N, 7.01.

3.3.4. 9-Allyl-2,3-dihydro-1H-pyrrolo[1,2-a]indole (9d). IR (neat): 1660, 1640, 1610 cm⁻¹. ¹H NMR (CDCl₃) δ : 2.55 (tt, 2H, $J=7.2$, 7.3 Hz), 2.92 (t, 2H, $J=7.3$ Hz), 3.45 (d, 2H, $J=5.6$ Hz), 3.99 (t, 2H, $J=7.2$ Hz), 5.00 (dd, 1H, $J=1.5$, 10.0 Hz), 5.10 (dd, 1H, $J=1.7$, 17.0 Hz), 6.02 (ddt, 1H, $J=17.0$, 10.0, 5.6 Hz), 7.03 (dd, 1H, $J=7.8$, 7.9 Hz), 7.09 (dd, 1H, $J=7.8$, 7.9 Hz), 7.18 (d, 1H, $J=7.8$ Hz), 7.49 (d, 1H, $J=7.9$ Hz). ¹³C NMR (CDCl₃) δ : 23.1, 27.6, 29.3, 43.3, 102.9, 109.1, 114.3, 118.4, 118.5, 119.9, 132.3, 132.4, 137.4, 141.5. HR-MS m/z : Calcd for C₁₄H₁₅N: 197.1203. Found: 197.1220.

3.3.5. 10-Allyl-6,7,8,9-tetrahydropyrido[1,2-a]indole (9e). ¹H NMR (CDCl₃) δ : 1.88–1.94 (m, 2H), 2.05–2.11 $(m, 2H), 2.89$ (t, $2H, J=6.3$ Hz), 3.47 (d, $2H, J=6.3$ Hz), 4.04 (t, 2H, $J=6.3$ Hz), 5.01 (dd, 1H, $J=1.0$, 10.1 Hz), 5.09 (dd, 1H, $J=1.0$, 17.1 Hz), 5.99 (tdd, 1H, $J=6.3$, 10.1, 17.1 Hz), 7.09 (t, 1H, $J=7.8$ Hz), 7.15 (t, 1H, $J=7.8$ Hz), 7.26 (d, 1H, $J=7.8$ Hz), 7.54 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 21.2, 22.5, 23.5, 28.5, 42.4, 107.0, 108.5, 114.2, 118.0, 119.1, 120.2, 127.9, 133.5, 136.0, 137.5. HR-MS m/z: Calcd for $C_{15}H_{17}N$: 211.1361. Found: 211.1353.

3.3.6. 11-Allyl-7,8,9,10-tetrahydro-6H-azepino[1,2-a] **indole (9f).** ^IH NMR (CDCl₃) δ : 1.67–1.73 (m, 2H), 1.73–1.79 (m, 2H), 1.81–1.87 (m, 2H), 2.85 (t, 2H, $J=$ 5.2 Hz), 3.48 (td, 2H, $J=1.7$, 6.3 Hz), 4.15 (t, 2H, $J=5.8$ Hz), 4.96 (qd, 1H, $J=1.5$, 11.4 Hz), 5.03 (qd, 1H, $J=1.5$, 17.2 Hz), 5.96 (tdd, 1H, $J=6.3$, 11.4, 17.2 Hz), 7.03 (t, 1H, $J=7.8$ Hz), 7.13 (dt, 1H, $J=1.5$, 7.8 Hz), 7.25 (d, 1H, $J=8.0$ Hz), 7.50 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) d: 25.5, 27.9, 28.8, 29.6, 31.2, 44.6, 107.7, 108.4, 114.1, 118.4, 118.5, 120.5, 127.6, 135.8, 138.1, 139.6. HR-MS m/z: Calcd for $C_{16}H_{19}N: 225.1517$. Found: 225.1526.

3.3.7. 2,3-Dihydro-1H-pyrrolo[1,2-a]indol-9-ylacetonitrile (9g). Mp 99–100 °C (hexane). IR (CHCl₃): 2430 cm⁻¹.²¹H NMR (CDCl₃) δ : 2.55–2.65 (m, 2H), 3.03 $(t, 2H, J=7.3 \text{ Hz})$, 3.73 (s, 2H), 4.00 (t, 2H, $J=6.8 \text{ Hz}$), 7.11 (dt, 1H, $J=1.0$, 7.8 Hz), 7.15 (dt, 1H, $J=1.0$, 7.8 Hz), 7.21 (d, 1H, $J=7.8$ Hz), 7.46 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl3) d: 13.5, 23.1, 27.5, 43.7, 93.4, 109.6, 117.5, 118.1, 119.3, 120.9, 130.9, 132.3, 142.5. MS m/z : 196 (M⁺). Anal. Calcd for $C_{13}H_{12}N_2$: C, 79.56; H, 6.16; N, 14.27. Found: C, 79.58; H, 6.25; N, 14.26.

3.3.8. 6,7,8,9-Tetrahydropyrido[1,2-a]indol-10-ylacetonitrile (9h). Mp $102-103$ °C (hexane). IR (CHCl₃): 2428 cm⁻¹. ¹H NMR (CDCl₃) δ : 1.91-1.97 (m, 2H), 2.06–2.12 (m, 2H), 2.94 (t, 2H, $J=6.3$ Hz), 3.74 (s, 3H), 4.04 (t, 2H, $J=6.3$ Hz), 7.16 (t, 1H, $J=7.8$ Hz), 7.19 (t, 1H, $J=7.8$ Hz), 7.28 (d, 1H, $J=8.0$ Hz), 7.55 (d, 1H, $J=$ 8.0 Hz). 13C NMR (CDCl3) d: 12.7, 20.7, 22.3, 23.1, 42.3, 97.5, 108.9, 117.1, 118.2, 120.1, 121.1, 126.6, 134.6, 135.9. MS m/z : 210 (M⁺). Anal. Calcd for C₁₄H₁₄N₂: C, 79.97; H, 6.71; N, 13.32. Found: C, 79.74; H, 6.74; N, 13.20.

3.3.9. 7,8,9,10-Tetrahydro-6H-azepino[1,2-a]indol-11-ylacetonitrile (9i). Mp $102-103^{\circ}$ C (hexane). IR (CHCl₃): 2430 cm⁻¹. ¹H NMR (CDCl₃) δ : 1.73–1.80 (m, 4H), $1.85-1.90$ (m, 2H), 2.90 (t, 2H, $J=5.1$ Hz), 3.78 (s, 3H), 4.17 (t, 2H, $J=5.1$ Hz), 7.13 (t, 1H, $J=7.8$ Hz), 7.20 (dt, 1H, $J=1.5, 7.8$ Hz), 7.28 (d, 1H, $J=8.0$ Hz), 7.55 (d, 1H, $J=$ 8.0 Hz). ¹³C NMR (CDCl₃) δ : 13.1, 25.6, 27.4, 29.3, 31.0, 44.8, 98.6, 108.9, 117.6, 118.5, 119.5, 121.4, 126.4, 135.8, 140.6. MS *m/z*: 224 (M⁺). Anal. Calcd for C₁₅H₁₆N₂: C, 80.32; H, 7.19; N, 12.49. Found: C, 80.22; H, 7.33; N, 12.35.

3.3.10. 9-Cyclohex-2-en-1-yl-2,3-dihydro-1H-pyrrolo- [1,2-a]indole (9j). Mp $53-54\,^{\circ}\text{C}$ (hexane). ¹H NMR (CDCl₃) δ : 1.62–1.71 (m, 1H), 1.72–1.83 (m, 2H9), 1.96–2.04 (m, 1H), 2.08–2.14 (m, 2H), 2.52–2.60 (m, 2H), 2.91–3.03 (m, 2H), 3.66–3.71 (m, 1H), 4.01 (t, 2H, $J=$ 7.0 Hz), 5.83 (s, 1H), 7.01 (t, 1H, $J=7.8$ Hz), 7.09 (t, 1H, $J=7.8$ Hz), 7.20 (d, 1H, $J=8.0$ Hz), 7.55 (d, 1H, $J=$ 8.0 Hz). ¹³C NMR (CDCl₃) δ : 21.5, 24.1, 25.2, 27.7, 30.6, 32.7, 43.1, 109.1, 109.5, 118.2, 118.7, 119.9, 127.0, 131.0, 131.6, 132.3, 140.9. MS m/z : 237 (M⁺). Anal. Calcd for $C_{17}H_{19}N+1/10H_2O$: C, 85.38; H, 8.09; N, 5.85. Found: C, 85.30; H, 8.22; N, 5.90.

3.3.11. rel-(9aR,10S)-10-Cyclohex-2-en-1-yl-9a-(9 oxa-10-borabicyclo[3.3.2]dec-10-yl)-6,7,8,9,9a,10-hexahydropyrido[1,2-a]indole (9k). Mp $209-210$ °C (hexane/ AcOEt). ¹H NMR (CDCl₃) δ : 0.74–0.83 (m, 1H), 1.30–2.10 $(m, 24H), 2.40$ (dd, 1H, $J=2.8, 5.1$ Hz), 2.91 (d, 1H, $J=$ 2.8 Hz), 3.51 (dt, 1H, $J=2.8$, 13.4 Hz), 3.60 (d, 1H, $J=$ 13.4 Hz), 4.80–4.86 (m, 1H), 5.63 (d, 1H, $J=10.3$ Hz), 5.72 (td, 1H, $J=1.8$, 10.3 Hz), 6.48 (d, 1H, $J=7.5$ Hz), 6.54 (t, 1H, $J=7.5$ Hz), 6.99 (d, 1H, $J=6.9$ Hz), 7.05 (t, 1H, $J=$ 8.0 Hz). ¹³C NMR (CDCl₃) δ : 21.1, 21.6, 22.1, 22.6, 23.6, 24.7, 24.8, 26.9, 27.1, 31.4, 33.9, 40.2, 43.9, 56.9, 67.6, 74.3, 108.2, 116.2, 126.2, 127.3, 128.7, 130.8, 131.8, 152.8. MS *m/z*: 388, 389 (M⁺). Anal. Calcd for $C_{26}H_{36}BNO+1/$ 4H2O: C, 79.28; H, 9.34; N, 3.55. Found: C, 79.03; H, 9.35; N, 3.53.

3.3.12. 11-Cyclohex-2-en-1-yl-7,8,9,10-tetrahydro-6Hazepino[1,2-a]indole (91). Mp $94-95$ °C (hexane). ¹H NMR (CDCl₃) δ: 1.62–1.94 (m, 10H), 2.09–2.25 (m, 2H), 2.83–2.93 (m, 2H), 3.63–3.70 (m, 1H), 4.09–4.19 (m, 2H), 5.79 (d, 1H, $J=10.3$ Hz), 5.85 (td, 1H, $J=2.3$, 9.7 Hz), 6.99 $(t, 1H, J=8.0 \text{ Hz})$, 7.10 $(t, 1H, J=8.0 \text{ Hz})$, 7.24 $(d, 1H, J=$ 8.4 Hz), 7.63 (d, 1H, J=7.8 Hz). ¹³C NMR (CDCl₃) δ : 23.0, 25.2, 25.4, 28.2, 29.5, 31.2, 31.7, 33.7, 44.4, 108.4, 114.0, 118.1, 119.4, 120.2, 126.9, 127.0, 132.5, 135.8, 139.0. MS m/z: 265 (M⁺). Anal. Calcd for C₁₉H₂₃N+1/10H₂O: C, 85.40; H, 8.75; N, 5.24. Found: 85.24; H, 8.72; N, 5.16.

3.3.13. Methyl $(2E)$ -4- $(2,3$ -dihydro-1H-pyrrolo[1,2-a]indol-9-yl)but-2-enoate $(9m)$. IR $(CHCl₃)$: 1710, 1648 cm^{-1} . ¹H NMR (CDCl₃) δ : 2.50–2.65 (m, 2H), 2.92 (t, 2H, $J=7.5$ Hz), 3.60 (d, 2H, $J=6.3$ Hz), 3.69 (s, 3H), 4.04 (t, 2H, $J=6.8$ Hz), 5.84 (td, 1H, $J=1.8$, 15.4 Hz), 7.05 (t, 1H, $J=7.8$ Hz), 7.10–7.19 (m, 2H), 7.21 (d, 1H, $J=$ 8.0 Hz), 7.43 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 23.3, 27.8, 27.9, 43.7, 51.4, 100.9, 109.5, 118.3, 118.9, 120.4, 120.9, 132.1, 132.6, 142.2, 148.1, 167.3. MS m/z: Calcd for $C_{16}H_{17}NO_2$: 255.1259. Found: 255.1258.

3.3.14. Methyl (2E)-4-(6,7,8,9-tetrahydropyrido[1,2-a] indol-10-yl)-but-2-enoate (9n). Mp 84–85 \degree C (hexane). IR (CHCl₃): 1710, 1654 cm⁻¹. ¹H NMR (CDCl₃) δ : 1.87-1.93 $(m, 2H), 2.03-2.11$ $(m, 2H), 2.83$ $(t, 2H, J=6.3$ Hz), 3.57 (dd, 2H, $J=1.7$, 6.3 Hz), 3.67 (s, 3H), 4.03 (t, 2H, $J=$ 6.3 Hz), 5.78 (td, 1H, $J=1.7$, 15.5 Hz), 7.08 (t, 1H, $J=$ 7.8 Hz), 7.14 (t, 1H, $J=7.8$ Hz), 7.25 (d, 1H, $J=8.0$ Hz), 7.43 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 21.1, 22.4, 23.4, 26.9, 42.3, 51.4, 104.8, 108.6, 117.7, 119.4, 120.5, 120.6, 127.6, 134.0, 136.0, 148.0, 167.3. MS m/z : 269 (M⁺). Anal. Calcd for $C_{17}H_{19}NO_2$: C, 75.80; H, 7.11; N, 5.20. Found: C, 75.71; H, 6.93; N, 5.11.

3.3.15. Methyl (2E)-4-(7,8,9,10-tetrahydro-6H-azepino- $[1,2-a]$ indol-11-yl)but-2-enoate (90). IR $(CHCl₃)$: 1708 cm^{-1} . ¹H NMR (CDCl₃) δ : 1.66–1.72 (m, 2H), $1.74-1.80$ (m, 2H), $1.83-1.90$ (m, 2H), 2.82 (t, 2H, $J=$ 5.2 Hz), 3.61 (dd, 2H, $J=1.7$, 6.3 Hz), 3.67 (s, 3H), 4.16 (t, $2H, J=4.6$ Hz), 5.75 (dt, 1H, $J=1.7$, 15.5 Hz), 7.05 (t, 1H, $J=8.0$ Hz), 7.11 (dt, 1H, $J=6.3$, 15.5 Hz), 7.17 (t, 1H, $J=$ 8.0 Hz), 7.25 (d, 1H, $J=8.0$ Hz), 7.42 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl₃) δ : 25.6, 27.3, 27.9, 29.5, 31.2, 42.0, 51.4, 105.6, 108.6, 118.1, 118.8, 120.7, 120.8, 127.4, 136.0, 140.2, 148.7, 167.4. MS m/z : Calcd for C₁₈H₂₁NO₂: 283.1572. Found: 283.1570.

3.3.16. (2E)-4-(2,3-Dihydro-1H-pyrrolo[1,2-a]indol-9 yl)but-2-en-1-ol (9p). IR (CHCl₃): 3612 cm⁻¹. ¹H NMR $(CDCl₃)$ δ : 2.55–2.65 (m, 2H), 2.93 (t, 2H, $J=7.5$ Hz), 3.46 $(d, 2H, J=6.9 \text{ Hz})$, 4.03 (t, 2H, $J=6.8 \text{ Hz}$), 4.09 (br s, 2H), $5.71-5.78$ (m, 1H), 5.90 (ttd, 1H, $J=1.2, 6.3, 15.5$ Hz), 7.03 $(dt, 1H, J=1.0, 7.8 Hz), 7.10 (dt, 1H, J=1.0, 7.8 Hz), 7.20$ (d, 1H, $J=8.0$ Hz), 7.48 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 23.3, 27.7, 43.4, 63.4, 103.1, 109.2, 118.5, 120.3, 129.0, 131.6, 132.2, 132.5, 141.5. HR-MS m/z: Calcd for $C_{15}H_{17}NO: 227.1310.$ Found: 227.1305.

3.3.17. (2E)-4-(6,7,8,9-Tetrahydro-1H-pyrido[1,2-a] indol-10-yl)but-2-en-1-ol (9q). IR (CHCl₃): 3612 cm⁻¹.
¹H NMP (CDCl) δ : 1.86, 1.93 (m, 2H) 2.03, 2.10 (m, 2H) ¹H NMR (CDCl₃) δ : 1.86–1.93 (m, 2H), 2.03–2.10 (m, 2H), 2.87 (t, 2H, $J=6.9$ Hz), 3.44 (d, 2H, $J=6.3$ Hz), 4.03 (t, 2H, $J=6.3$ Hz), 4.07 (br s, 2H), 5.65–5.74 (m, 1H), 5.80–5.88 $(m, 1H)$, 7.07 (t, 1H, $J=8.0$ Hz), 7.13 (t, 1H, $J=8.0$ Hz), 7.24 (d, 1H, $J=8.0$ Hz), 7.44 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl3) d: 21.2, 22.5, 23.5, 26.9, 63.8, 107.1, 108.6, 117.9, 119.2, 120.3, 127.8, 128.8, 131.9, 133.4, 136.0. HR-MS m/z: Calcd for $C_{16}H_{19}NO: 241.1466$. Found: 241.1466.

3.3.18. (2E)-4-(7,8,9,10-Tetrahydro-6H-azepino[1,2-a] **indol-11-yl)but-2-en-1-ol** (9r). IR (CHCl₃): 3612 cm^{-1} .
¹H NMP (CDCl) λ : 1.65, 1.01 (m, 6H) 2.85 (t, 2H, I ¹H NMR (CDCl₃) δ : 1.65–1.91 (m, 6H), 2.85 (t, 2H, J= 5.7 Hz), 3.48 (d, 2H, $J=5.7$ Hz), 4.06 (d, 2H, $J=6.3$ Hz), 4.15 (t, 2H, $J=4.6$ Hz), 5.64–5.71 (m, 1H), 5.84 (td, 1H, $J=5.7$, 15.1 Hz), 7.04 (t, 1H, $J=7.8$ Hz), 7.14 (t, 1H, $J=7.8$ Hz), 7.25 (d, 1H, $J=8.0$ Hz), 7.49 (d, 1H, $J=$ 8.0 Hz). 13C NMR (CDCl3) d: 25.6, 27.2, 28.0, 29.6, 31.2, 44.6, 63.6, 107.9, 108.5, 118.4, 118.6, 120.6, 127.5, 128.7, 132.5, 135.9, 139.6. HR-MS m/z : Calcd for C₁₇H₁₂NO: 255.1623. Found: 255.1612.

3.3.19. rel-(1R,4S)-4-(2,3-Dihydro-1H-pyrrolo[1,2-a] indol-9-yl)cyclohex-2-en-1-yl acetate (9s). Mp $101-02$ °C (hexane/AcOEt). IR (CHCl₃): 1720 cm^{-1} . ¹H NMR $(CDCl_3)$ δ : 1.86–1.94 (m, 1H), 1.94–2.00 (m, 1H), 2.09 (s, $3H$), 2.55–2.65 (m, 2H), 2.99 (t, 2H, $J=7.3$ Hz), 3.60–3.66 $(m, 2H), 4.02$ (t, $2H, J=7.3$ Hz), $5.30-5.36$ (m, 1H), 5.88 (ddd, 1H, $J=2.4$, 4.0, 10.0 Hz), 6.10 (dd, 1H, $J=2.4$, 10.0 Hz), 7.04 (t, 1H, $J=7.8$ Hz), 7.11 (t, 1H, $J=7.8$ Hz), 7.21 (d, 1H, $J=8.0$ Hz), 7.54 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 21.4, 24.2, 26.2, 27.4, 27.8, 32.9, 43.2, 67.5, 107.9, 109.3, 118.5, 118.6, 120.2, 125.1, 131.5, 132.5, 136.8, 141.1, 170.8. MS m/z : 295 (M⁺). Anal. Calcd for $C_{19}H_{21}NO_2$: C, 77.26; H, 7.17; N, 4.74. Found: C, 77.02; H, 7.12; N, 4.49.

3.3.20. 9-(Tetrahydrofuran-2-yl)-2,3-dihydro-1Hpyrrolo[1,2-a]indole (9t). Mp $72-73$ °C (hexane/AcOEt). ¹H NMR (CDCl₃) δ : 2.00–2.20 (m, 3H), 2.22–2.30 (m, 1H), 2.56–2.62 (m, 2H), 2.99–3.11 (m, 2H), 3.88–3.94 (m, 1H), 4.02 (t, 2H, $J=6.9$ Hz), 4.06–4.12 (m, 1H), 5.16 (t, 1H, $J=$ 7.4 Hz), 7.05 (t, 1H, $J=8.0$ Hz), 7.11 (t, 1H, $J=8.0$ Hz), 7.20 (d, 1H, $J=8.0$ Hz), 7.59 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 24.2, 26.6, 27.4, 43.5, 68.0, 75.4, 106.7, 109.5, 119.0, 119.4, 120.4, 131.1, 132.9, 141.8. MS m/z : 227 (M⁺). Anal. Calcd for $C_{15}H_{17}NO: C$, 79.26; H, 7.54; N, 6.16. Found: C, 79.05; H, 7.68; N, 5.99.

3.3.21. 3- $(1H$ -Indol-1-yl)propan-1-ol $(10a)$. IR $(CHCl₃)$: 3616 cm⁻¹. ¹H NMR (CDCl₃) δ : 1.56 (br s, 1H), 2.03–2.10 (m, 2H), 3.58 (t, 2H, $J=5.7$ Hz), 4.28 (t, 2H, $J=6.8$ Hz), 6.51 (d, 1H, $J=2.9$ Hz), 7.09–7.15 (m, 2H), 7.22 (t, 1H, $J=$ 7.8 Hz), 7.39 (d, 1H, $J=8.0$ Hz), 7.64 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 32.7, 42.7, 59.6, 101.2, 109.4, 119.4, 121.1, 121.5, 128.0, 128.7, 136.1. HR-MS m/z: Calcd for $C_{11}H_{13}NO: 175.0997.$ Found: 175.0982.

3.3.22. 3-(5-Nitro-1H-indol-1-yl)propan-1-ol (10b). Mp 85 °C (hexane/AcOEt). IR (CHCl₃): 3628 cm^{-1} . ¹H NMR $(CDCl_3)$ δ : 1.57 (br s, 1H), 2.03–2.12 (m, 2H), 3.61 (t, 2H, $J=6.3$ Hz), 4.34 (t, 2H, $J=6.9$ Hz), 6.68 (d, 1H, $J=$ 2.3 Hz), 7.28 (d, 1H, $J=2.3$ Hz), 7.41 (d, 1H, $J=9.0$ Hz), 8.10 (dd, 1H, $J=1.7$, 8.6 Hz), 8.58 (d, 1H, $J=1.7$ Hz). ¹³C NMR (CDCl₃) δ: 32.6, 43.1, 59.0, 104.1, 109.3, 117.3, 118.3, 127.7, 131.2, 139.0, 141.6. MS m/z : 220 (M⁺). Anal. Calcd for $C_{11}H_{12}N_2O_3$: C, 59.99; H, 5.49; N, 12.72. Found: C, 59.76; H, 5.51; N, 12.89.

3.3.23. 4- $(1H$ -Indol-1-yl)butan-1-ol $(10c)$. IR $(CHCl₃)$: 3624 cm^{-1} . ¹H NMR (CDCl₃) δ : 1.50–1.55 (m, 2H), 1.68 (br s, 1H), $1.87-1.94$ (m, 2H), 3.59 (t, 2H, $J=6.3$ Hz), 4.15 $(t, 2H, J=6.8 \text{ Hz})$, 6.55 (d, 2H, $J=2.9 \text{ Hz}$), 7.09–7.14 (m, 2H), 7.22 (t, 1H, $J=8.0$ Hz), 7.36 (d, 1H, $J=8.0$ Hz), 7.64 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl₃) δ : 26.8, 30.1, 46.2, 62.3, 101.1, 109.5, 119.4, 121.1, 121.5, 127.9, 128.7, 136.0. HR-MS m/z : Calcd for C₁₂H₁₅NO: 189.1154. Found: 189.1166.

3.3.24. 5- $(1H$ -Indol-1-yl)pentan-1-ol $(10d)$. IR $(CHCl₃)$: 3620 cm^{-1} . ¹H NMR (CDCl₃) δ : 1.35–1.40 (m, 2H), 1.53–1.60 (m, 2H), 1.83–1.90 (m, 2H), 3.60 (t, 2H, $J=$ 6.3 Hz), 4.13 (t, 2H, $J=6.8$ Hz), 6.49 (d, 1H, $J=2.9$ Hz), 7.08–7.12 (m, 2H), 7.21 (t, 1H, $J=8.0$ Hz), 7.34 (d, 1H, $J=8.0$ Hz), 7.63 (d, 1H, $J=7.8$ Hz). ¹³C NMR (CDCl₃) δ : 23.4, 30.2, 32.4, 46.4, 62.7, 101.0, 109.4, 119.3, 121.1, 121.4, 127.9, 128.7, 136.0. HR-MS m/z: Calcd for C13H17NO: 203.1310. Found: 203.1295.

3.4. General procedure for the preparation of 13, 14 and 17

The reaction using 11 was carried out according to the procedure for the prepatation of 8. Hydroboration of 11b with 9-BBN was effected under reflux for 2 h.

3.4.1. 2-Methyl-2,3-dihydro-1H-pyrrolo[1,2-a]indole (13a). Mp 58-59 °C (hexane/AcOEt). ¹H NMR (CDCl₃) δ : 1.29 (d, 3H, J=6.9 Hz), 2.63 (dd, 1H, J=6.9, 15.9 Hz), $3.04-3.14$ (m, 1H), 3.19 (dd, 1H, $J=8.0$, 15.9 Hz), 3.62 (dd, 1H, $J=6.3$, 9.7 Hz), 4.22 (dd, 1H, $J=7.4$, 9.7 Hz), 6.15 (s, 1H), 7.05 (dt, 1H, $J=1.0$, 7.8 Hz), 7.11 (td, 1H, $J=1.0$, 7.8 Hz), 7.21 (d, 1H, 8.0 Hz), 7.54 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ: 19.7, 33.0, 37.3, 50.9, 92.6, 109.3, 119.1, 120.2, 120.3, 132.8, 132.9, 144.2. MS m/z : 171 (M⁺). Anal. Calcd for $C_{12}H_{13}N$: C, 84.17; H, 7.65; N, 8.18. Found: C, 84.15; H, 7.64; N, 8.16.

3.4.2. 2-Phenyl-2,3-dihydro-1H-pyrrolo[1,2-a]indole (13b). ¹H NMR (CDCl₃) δ : 3.17 (dd, 1H, J=7.5, 15.9 Hz), 3.50 (dd, 1H, $J=8.0$, 15.9 Hz), 4.09 (dd, 1H, $J=6.8$, 9.9 Hz), 4.15–4.22 (m, 1H), 4.51 (dd, 1H, $J=8.1$, 9.9 Hz), 6.27 (s, 1H), 7.13 (t, 1H, $J=7.8$ Hz), 7.18 (t, 1H, $J=7.8$ Hz), $7.26-7.33$ (m, 4H), $7.35-7.40$ (m, 2H), 7.63 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 33.3, 47.9, 51.1, 93.0, 109.5, 119.4, 120.5, 120.6, 127.1, 127.2, 129.0, 132.9, 133.2, 142.9, 143.6. HR-MS m/z : Calcd for C₁₇H₁₅N: 233.1204. Found: 233.1210.

3.4.3. 3-(1H-Indol-1-yl)-2-methylpropan-1-ol (14a). IR (CHCl₃): 3624 cm⁻¹. ¹H NMR (CDCl₃) δ : 0.97 (d, 3H, J= 6.9 Hz), 1.37 (br s, 1H), 2.20–2.29 (m, 1H), 3.49 (d, 2H, $J=$ 4.0 Hz), 3.99 (dd, 1H, $J=6.9$, 14.3 Hz), 4.24 (dd, 1H, $J=$ 6.9, 14.3 Hz), 7.17–7.12 (m, 2H), 7.20 (t, 1H, $J=8.0$ Hz), 7.38 (d, 1H, $J=8.0$ Hz), 7.63 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 15.0, 36.9, 49.0, 65.1, 101.2, 109.6, 119.3, 121.0, 121.5, 128.6, 136.4. HR-MS m/z : Calcd for C₁₂H₁₅NO: 189.1154. Found: 189.1153.

3.4.4. $3-(1H\text{-Indol-1-vl})-2\text{-phenylindole}$ (14b). IR (CHCl₃): 3624 cm^{-1} . ¹H NMR (CDCl₃) δ : 1.42 (br s, 1H), $3.31-3.38$ (m, 1H), 3.78 (br s, 2H), 4.33 (dd, 1H, $J=$ 6.9, 14.4 Hz), 4.59 (dd, 1H, $J=8.0$, 14.4 Hz), 6.42 (d, 1H, $J=3.5$ Hz), 6.92 (d, 1H, $J=3.5$ Hz), 7.10 (t, 1H, $J=$ 7.8 Hz), 7.18–7.23 (m, 3H), 7.25–7.31 (m, 1H), 7.31–7.39 (m, 3H), 7.62 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl₃) δ : 48.3, 48.7, 64.1, 101.3, 109.5, 119.5, 121.1, 121.6, 127.5, 128.1, 128.5, 128.7, 129.0, 136.2, 139.9. HR-MS m/z: Calcd for C17H17NO: 251.1301. Found: 231.1304.

3.4.5. 1-Allyl-1,3-dihydro-2H-indol-2-one (17). IR (neat): 1708 cm⁻¹.¹H NMR (CDCl₃) δ : 3.55 (s, 2H), 4.34 (dt, 2H, $J=5.1, 1.5$ Hz), 5.21 (dd, 1H, $J=1.0$, 10.3 Hz), 5.23 (dd, $1H, J=1.0, 17.1$ Hz), 5.84 (tdd, $1H, J=5.1, 10.3, 17.1$ Hz), 6.81 (d, 1H, $J=7.8$ Hz), 7.02 (dt, 1H, $J=1.0$, 7.8 Hz), 7.23 (t, 1H, $J=7.8$ Hz), 7.24 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 35.7, 42.2, 108.9, 117.5, 122.3, 124.3, 124.4,

127.7, 131.4, 144.3, 174.7. HR-MS m/z: Calcd for $C_{11}H_{11}NO: 173.0840.$ Found: 173.0822.

3.5. Procedure for the preparation of 9a, 20, 10e and 21 from 18

Conversion of 18a to 9a (40%) and 10a (13%) was carried out according to the procedure for the preparation of 8.

Reaction using 18b was effected as follows: to a solution of 18b (430 mg, 2 mmol) in THF (10 mL), 9-BBN (0.5 M solution in THF, 2.2 mmol) was added at room temperature under an argon atmosphere, and the mixture was stirred for 1.5 h. Sodium methoxide (118 mg, 2.2 mmol) was added to the mixture, and after stirring for 30 min, LDA (4.4 mmol) was added to the mixture at -78 °C. After stirring for 1 h, the mixture was gradually raised to room temperature, and stirred for 4 h. To the reaction mixture, 20% NaOH (10 mL) and 30% $H₂O₂$ (2 mL) were added under ice-cooling and the whole was stirred for 30 min. The mixture was diluted with ethyl acetate (100 mL) , washed with brine, and dried over MgSO₄. The solvent was removed, and the residue was separated by MPLC to give 20 (69 mg, 16%) (with hexane/AcOEt = 50:1) and 21 $(47 \text{ mg}, 10\%)$ (with hexane/AcOEt = 1:1).

3.5.1. 3-(1H-3-Methylindol-1-yl)propan-1-ol (10e). IR (CHCl₃): 3624 cm⁻¹.¹H NMR (CDCl₃) δ : 2.01–2.07 (m, 2H), 2.32 (s, 3H), 3.58–3.63 (m, 2H), 4.21 (t, 2H, $J=$ 6.3 Hz), 6.88 (s, 1H), 7.09 (t, 1H, $J=7.8$ Hz), 7.19 (t, 1H, $J=7.8$ Hz), 7.33 (d, 1H, $J=8.0$ Hz), 7.56 (d, 1H, $J=$ 8.0 Hz). ¹³C NMR (CDCl₃) δ : 9.6, 32.8, 42.3, 59.7, 109.1, 110.4, 118.6, 119.1, 121.4, 125.6, 128.8, 136.4. HR-MS m/z: Calcd for $C_{12}H_{15}NO: 189.1154$. Found: 189.1142.

3.5.2. Methyl 2,3-dihydro-1H-pyrrolo[1,2-a]indole-9 carboxylate (20). Mp $92-93$ °C (hexane). IR (neat): 1686 cm^{-1} . ¹H NMR (CDCl₃) δ : 2.61–2.70 (m, 2H), 3.29 (t, 2H, $J=7.5$ Hz), 3.89 (s, 3H), 4.11 (t, 2H, $J=5.8$ Hz), 7.18–7.28 (m, 3H), 8.09 (d, 1H, $J=8.0$ Hz). ¹³C NMR (CDCl3) d: 26.2, 26.7, 44.5, 50.8, 99.5, 109.9, 121.5, 121.7, 121.8, 130.9, 132.7, 152.9, 166.0. HR-MS m/z : 215 (M⁺). Anal. Calcd for C₁₃H₁₃NO₂: C, 72.54; H, 6.09; N, 6.51. Found: 72.53; H, 6.11; N, 6.46.

3.5.3. Methyl 1-(3-hydroxypropyl)-1H-indole-3 carboxylate (21). IR (neat): 1690 cm^{-1} . ¹H NMR (CDCl3) d: 1.65 (br s, 1H), 2.03–2.11 (m, 2H), 3.60 (t, $2H, J=5.8$ Hz), 3.90 (s, 3H), 4.32 (t, 2H, $J=6.3$ Hz), 7.25– 7.29 (m, 2H), 7.39–7.42 (m, 1H), 8.15–8.19 (m, 1H). 13C NMR (CDCl₃) δ: 32.1, 43.2, 51.1, 59.0, 107.1, 110.0, 121.8, 121.9, 122.8, 126.7, 134.6, 136.6, 165.7. HR-MS m/z: C13H15NO3: 233.1051. Found: 233.1042.

3.5.4. rel-(9R,9aS)-9-Allyl-9a-(9-borabicyclo[3.3.1]non-9-yl)-9-methyl-2,3,9,9a-tetrahydro-1H-pyrrolo[1,2-a] indole (22). To a solution of 18a (342 mg, 2 mmol) in THF (10 mL) , 9-BBN $(0.5 \text{ M}$ solution in THF, 2.2 mmol) was added at room temperature under an argon atmosphere, and the mixture was stirred for 1.5 h. Sodium methoxide (118 mg, 2.2 mmol) was added to the mixture, and after stirring for 30 min, TMEDA (0.6 mL, 4.4 mmol) and tert-BuLi (1.5 M solution in pentane, 4.4 mmol) were added to the mixture at -20 °C. After stirring for 2 h, the

mixture was gradually raised to room temperature, and stirred for 4 h. To the reaction mixture, allyl bromide (605 mg, 5 mmol) was added and the mixture was stirred at room temperature overnight. The mixture was diluted with ethyl acetate (100 mL), washed with brine, and dried over $MgSO₄$. The solvent was removed, and the residue was separated by MPLC (hexane/AcOEt=100:1) to give 22 (232 mg, 35%).

Mp 127–128 °C (hexane/AcOEt). ¹H NMR (CDCl₃) δ : 0.03 (br s, 1H), 0.71 (br s, 1H), 1.22 (s, 3H), 1.15–1.35 $(m, 1H), 1.38-1.88$ $(m, 13H), 2.03$ $(dd, 1H, J=7.4,$ 12.6 Hz), 2.10–2.20 (m, 1H), 2.29 (dd, 1H, $J=9.7$, 13.2 Hz), 2.64 (td, 1H, $J=2.3$, 13.2 Hz), 3.06 (dt, 1H, $J=6.8$, 11.4 Hz), 3.35 (t, 1H, $J=9.7$ Hz), 5.12 (d, 1H, $J=$ 10.1 Hz), 5.16 (d, 1H, $J=17.2$ Hz), 5.96 (dtd, 1H, $J=4.6$, 10.1, 17.2 Hz), 6.98 (dd, 1H, $J=1.7$, 8.3 Hz), 7.05 (dd, 1H, $J=1.7$, 7.8 Hz), 7.10–7.17 (m, 2H). ¹³C NMR (CDCl3) d: 21.0, 23.4, 24.1, 26.6, 26.8, 31.6, 31.9, 32.7, 33.7, 43.1, 48.7, 52.1, 117.2, 117.9, 123.1, 125.7, 126.8, 136.7, 146.6, 148.2. MS m/z : 332, 333 (M⁺). Anal. Calcd for $C_{23}H_{32}BN$: C, 82.87; H, 9.67; N, 4.20. Found: C, 82.74; H, 9.77; N, 4.16.

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Asymmetric Diels–Alder reactions with hydrogen bonding heterogeneous catalysts and mechanistic studies on the reversal of enantioselectivity

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Abstract—Chiral bis(oxazoline) complexes of Cu(II), $Zn(II)$ and Mg(II) have been immobilized on silica support via hydrogen-bonding interactions. Up to 93% ee is obtained in the Diels–Alder reaction between 3-((E)-2-butenoyl)-1,3-oxazolin-2-one and cyclopentadiene at room temperature with the heterogeneous bis(oxazoline) complexes, and the catalysts can be recycled without losing enantioselectivity. Experimental and theoretical studies show that the reversal of the absolute product configuration upon immobilization of the PhBOX-Cu(II) catalyst is triggered by the anion dissociation from Cu(II) onto the surface of the support. $© 2005 Elsevier Ltd. All rights reserved.$

1. Introduction

Enantioselective reactions catalyzed by chiral Lewis acid complexes are of great importance for the production of enantiopure pharmaceuticals and chemicals.^{[1](#page-193-0)} Among various chiral Lewis acid catalysts, those containing the chiral bis(oxazoline) (BOX) ligands have shown many applications in the last decade or so. $1,2$ Particularly, excellent enantioselectivities have been obtained in DA reactions using this type of the complexes.^{2,3} However, relatively large amounts of the chiral catalysts (1–10 mol%) are generally required, which makes recovery and recycling of the catalyst necessary. A number of strategies have therefore been designed and employed to immobilize and recycle the complexes of BOX ligands; $⁴$ these include</sup> covalent bonding to organic polymers and inorganic supports,^{[5,6](#page-193-0)} non-covalent immobilization by the interaction between cationic BOX-based metal complexes and anionic supports, $7,8$ and the use of solvents of special properties such as ionic liquids.^{[9](#page-194-0)}

Non-covalent immobilization is usually a convenient and also an industrially relevant method.[10](#page-194-0) However, few examples of non-covalently immobilized BOX systems have been reported or known to be effective for the DA reactions at present. Mayoral et al. exchanged the BOX- $M(II)$ (M = Cu, Mg, Zn) complexes onto laponite clays and nafion–silica nanocomposites for a benchmark DA reaction; but the ee was low (11%) .^{[8a](#page-194-0)} Hutchings et al. exchanged zeolite and mesoporous materials (MCM-41, Al-SBA-15, MSU-2) with $Cu(OAc)$ to obtain $Cu(II)$ -exchanged materials, then modified the exchanged materials with chiral BOX ligands for a hetero-DA reaction. The resulting PhBOX-CuH-zeolite Y catalyst gave a higher enantioselectivity (41% ee) compared to the homogeneous analogue (20% ee) although the activity was relatively low.^{[8b](#page-194-0)} More recently, the immobilization of homogeneous catalysts by hydrogen bonding has been reported.^{[11](#page-194-0)} Parallel to our research in immobilizing BOX complexes by hydrogen bonding for the DA reactions, Klein Gebbink and co-workers reported the same strategy for similar reactions.[12](#page-194-0) A surprising observation from these studies is that the configuration of the product changed on going from the homogeneous to the heterogeneous system. This is of both fundamental and practical significance, as it indicates that immobilization alters the active catalytic species, and both enantiomers of a product may be accessible by choosing a suitable support. However, the origin of the

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observed reversal is not yet clear, although it may be attributed to the interactions of the immobilized complexes with support surface. $8b,12$ Herein we describe our results of the DA reaction of dienophile 5 with diene 6 using hydrogen bond-immobilized BOX-M(II) catalysts [Eq. 1] and the insight we gained both experimentally and computationally into the reversal of product configuration.

The immobilization of the homogeneous catalysts H1–H4 onto silica was assumed to derive from hydrogen bonding interactions between the triflate anions and surface silanol groups (Fig. 1). Direct evidence for the hydrogen bonding is obtained from IR studies. Before adsorption, the support displays a sharp peak at 3740 cm⁻¹, which can be ascribed

2. Results and discussion

2.1. Preparation and characterization of the catalysts

The homogeneous catalysts H1–H4 were prepared in dichloromethane using metal triflates and 1 equiv of a BOX ligand. The corresponding supported catalysts S1–S4 were prepared by the adsorption of complexes H1–H4 to the surface of the silica (Fig. 1). The loading of the metal in the resulting heterogeneous catalysts is 1.0–1.1 wt% for copper and zinc, but only 0.03 wt% for magnesium. Anhydrous dichloromethane was used as the solvent for the preparation of the heterogeneous catalysts because of its dissolubility for BOX catalysts and aprotic nature (protic solvents would interfere with the adsorption).^{[13](#page-194-0)} Before the adsorption, the support was heated at $300\degree C$ in order to remove the physisorbed water, which would otherwise complicate the interpretation of the results (vide infra).

H1: $M = Cu, R = tBu$	S ₁	
$H2: M = Cu, R = Ph$	S ₂	
H3: $M = Zn$, $R = Ph$	S ₃	
H4: $M = Mg$, $R = Ph$	S4	

Figure 1. The precatalysts BOX-M(II) H1–H4 and their immobilization by hydrogen-bonding interaction.

to the stretching frequency of isolated silanols (Fig. 2a).^{[14](#page-194-0)} Upon adsorption of the homogeneous catalysts, the intensity of this IR band decreases considerably and only a shoulder at 3651 cm⁻¹ is observed. In addition, the bending vibration of the Si–O in silanols characterized by the band at 975 cm⁻¹ is red shifted to 946 cm⁻¹ and becomes broad (Fig. 2b).^{[14](#page-194-0)} Pimentel et al. showed that hydrogen bonding results in red-shifts in the –OH stretching vibration frequency and the half-width of the band is broadened.^{[13](#page-194-0)} This has been confirmed by others.^{[11a,15](#page-194-0)} Support for the integrity of the BOX structures in S1–S4 comes from the IR spectra [\(Fig. 3\)](#page-189-0).

Figure 2. FTIR spectra of the stretching and the bending frequencies of the hydroxyl groups on silica before and after the immobilization of H2: (a) the stretching frequencies of the hydroxyl groups; (b) the bending frequencies of the hydroxyl groups.

A rather intense band at \sim 1630 cm⁻¹ is observed, which corresponds to the $C=N$ bond stretching and is similar to that of the homogeneous complexes $(\sim 1633 \text{ cm}^{-1})$. We can therefore conclude that upon adsorption of the triflates complexes H1–H4 to silica, hydrogen-bonding interactions take place between silanols and sulfonate oxygen atoms, and the BOX-M(II) cations are immobilized primarily via ionic interaction.

Figure 3. FTIR spectra of the supported catalysts: t BuBOX-Cu(II)-SiO₂ (S1), PhBOX-Cu(II)-SiO₂ (S2), PhBOX-Zn(II)-SiO₂ (S3), and PhBOX- $Mg(II)$ -SiO₂ (S4).

2.2. Homogeneous versus heterogeneous DA reactions

For homogeneous DA reactions, better enantioselectivities are usually obtained at lower temperatures. However, practical applications of heterogeneous catalysts often need relatively high temperatures. With this in mind, the frequently used homogeneous catalysts H1 and H2 were initially immobilized for the DA reaction of dienophile 5 with diene **6** at room temperature [Eq. 1]. Table 1 summarizes the results obtained with **S1**, and for comparison those obtained with H1 in homogeneous reactions are also included. As we can see, the supported catalyst displays a lower activity and enantioselectivity compared to the homogeneous analogue in dichloromethane (entries 1 and 2). This is at least partially due to leaching of copper cations into the solution, which contained 79.8 ppm copper at the time when the reaction was terminated. Nevertheless,

the percentage of leaching was only 0.8%. A homogeneous experiment was carried out in which 0.8% amount of the catalyst was added and no more than 15% of conversion was observed, indicating a heterogeneous process. In contrast, an excellent enantioselectivity and enhanced activity were observed with S1 when the less polar toluene was used as the solvent (entries 3 and 4) in which the copper leaching is reduced to 4.9 ppm. The introduction of additional substrate to the filtrate before separating the product resulted in no obvious reaction, indicating that the DA reaction we observed is heterogeneous. However, a decreased activity was observed when the heterogeneous catalyst was reused (entries 5 and 6). Based on a study by Evans et al., who showed that the hydration of H1 brought about a detrimental effect on its activity and 3 Å molecular sieves proved effective in reactivating the catalyst, $3c$ we deduced that the activity decrease could arise from the effects of moisture or copper leaching in the subsequent reactions. 3 Å molecular sieves were therefore added to the heterogeneous system and we were delighted to find that S1 could be reused. After 3 runs, the enantioselectivity remained unchanged, although the activity decreased slightly (entries 7–9). The endo/exo ratios of H1 and S1 are comparable in both solvents.

The phenyl-substituted BOX ligand is known to be less enantioselective in Cu(II) catalyzed DA reactions than its t Bu analogue. This is also true with $S2$, as seen from [Table 2](#page-190-0). However, S2 provides higher enantioselectivities than H2 in all three solvents examined ([Table 2](#page-190-0), entries 1–4). This might be attributed to geometric constrains imposed on the PhBOX-Cu(II) catalyst by the silica surface. $8b,11b,c$ It may also arise from the reaction occurring at the liquid–solid interface, as the enantioselectivities of related reactions are known to be solvent-dependent.^{[3c,16](#page-193-0)} The lower ee values observed with $H2$ in toluene (3%) and ether (6%) may be also resulted from the lower solubility of the complex in the two solvents. In the case of S1, better enantioselectivity is obtained when toluene is used as the solvent in the heterogeneous catalysis ([Table 2](#page-190-0), entry 4). In contrast to S1, the phenyl-substituted S2 affords product with the configuration opposite to that obtained with its counterpart H2. This is interesting, as it suggests that the DA reaction with S2 is a heterogeneous process and more importantly, the active catalytic species are different on going from the homogeneous solution to the solid surface (vide infra).

Table 1. Homogeneous versus heterogeneous DA reactions catalyzed by the catalysts H1 and S1, respectively^a

Entry	Catalyst	Solvent	Time (h)	Conv. $(\%)^b$	endolexo ^c	ee $(\%)^c$	Config. d
	H1	CH_2Cl_2		$> 97^e$	89:11	90	
	S ₁	CH_2Cl_2		43	89:11	85	
	H1	Toluene		70	92:8	88	
4	S ₁	Toluene		83	91:9	93	
	S1(1st)	Toluene	22	98	89:11	91	
6	S1(2nd)	Toluene	22	76	92:8	92	
	S1(1st)	Toluene	22	98	90:10	91	
8 ^f	S1(2nd)	Toluene	22	92	92:8	91	
qt	S1(3rd)	Toluene	22	83	90:10	91	

^a Reactions were performed at room temperature with the ratio of catalyst/substrate=1/10. b The conversion was determined by ¹H NMR (400 MHz).

^c The *endolexo* ratio and the ee value of the *endo* isomer were analyzed by chiral HPLC. ^d The configuration of the product was confirmed by comparing with the literature.^{[3c](#page-193-0)}

 e° There was no peak of the substrate in the 1 H NMR spectrum.

 \hat{H} The reaction was carried out in the presence of 3 Å molecular sieves.

Entry	Catalyst	Solvent	Conv. $(\%)$	endolexo	ee $(\%)$	Config.	
	H2	CH_2Cl_2	100	89:11	15		
	S ₂	CH_2Cl_2	100	83:17			
	S ₂	Et ₂ O	100	90:10			
4	S2(1st)	Toluene	100	83:17	42		
	S2(2nd)	Toluene	100	82:18	46	ĸ	
6.	S2(3rd)	Toluene	98	84:16	40		
	$S2^b$	Toluene	100	79:21	21		
8	$S2^c$	Toluene	100	83:17	36		

Table 2. Homogeneous versus heterogeneous DA reactions catalyzed by the catalysts H2 and S2, respectively^a

^a The reaction time was 22 h under the conditions the same as those given in [Table 1.](#page-189-0)
^b The silica support was calcinated at 550 °C for 4 h.
^c The silica was firstly calcinated at 550 °C for 4 h, then refluxed in 1

We also attempted the reuse of **S2**. Following the first run of the DA reaction in toluene, the catalyst was firstly recovered by centrifugation and then dried for the next run. However, the activity and enantioselectivity of the catalyst decreased significantly. We have tried to improve reaction by working-up under an argon atmosphere. No significant change in selectivity is found when S2 is reused this way in toluene (Table 2, entries 4–6).

Previous results show that covering the free silanol groups on the silica increases the enantioselectivity in the DA reaction, 17 revealing the importance of the support effect. In our studies, different results are obtained. The silica involved was calcinated at 300° C for 3 h to remove the physisorbed water. However, when the silica was calcinated at 550 °C for 4 h (corresponding to partial dehydroxylation thus less silanol groups), 14 a significant decrease in ee was observed (Table 2, entries 4 and 7). Interestingly, when the silica calcinated at 550 \degree C was refluxed with 1 M HNO₃ for 2 h and washed with deionized water until neutral, the resulting catalyst gave an significant improved ee value (Table 2, entry 8). By treating with the $HNO₃$ for 2 h, the dehydroxylated surface of silica would be partially rehydroxylated and the silanol groups regenerated,^{[14](#page-194-0)} and hence the ee was improved again (Table 2, entries 7 and 8). These observations suggest that the change of the silanol groups on the surface of the support can influence the enantioselectivity. This is not surprising, as the closely related DA reaction of ethyl glyoxylate with 1,3-cyclohexdiene catalyzed by H2 has given ees depending on solvent dielectric constants[.16a](#page-194-0) The calcination of the support alters the density of surface silanol groups and so may affect their interaction with or the 'solvation' of the BOX complex.

2.3. Mechanistic studies on the reversal of enantioselectivity

Being interested in the observed reversal of the enantioselectivity (Table 2), we decided to undertake further studies. As aforementioned, the recent reports from the groups of Hutchings and Klein Gebbink have revealed similar observations.^{8b,12} However the cause for this reversal at the molecular level is still to be delineated. Such reversals in product configuration have also been noted in related homogeneous DA reactions on going from (S) -tBuBOX-Cu (II) to (S) -PhBOX-Cu(II) complexes.^{3d,18a,19} This was a result from the BOX-Cu(II)-dienophile complexes adopting a square planar vs. a tetrahedral geometry or from π -stabilization involving the PhBOX ligand. More recently, Jørgensen has demonstrated that, a number of factors such as solvents with different dielectric constants, may have subtle influence on the geometries of the BOX-Cu(II)-dienophile intermediates.^{16a}

For homogeneous asymmetric DA reactions, solvents, counterions and additives, for example, achiral ancillary ligands and molecular sieves, often influence the enantioface selection.^{[16a,18,20a,21](#page-194-0)} The asymmetric induction with BOX-M(II) catalysts is also dependent on the coordination geometry of the metal center, $3c,18,20$ with square planar and octahedral coordination favoring α -Si face addition while the tetrahedral arrangement favoring α -Re face reaction (Fig. 4). A distorted square pyramidal geometry is expected to give the same face selection as a square planar or octahedral species.

Figure 4. Effect of the BOX-M(II)-dienophile geometries on asymmetric induction: (a) planar, (b) tetrahedral, and (c) octahedral. M: metal, R: substituted group on the ring of BOX, L: coordinated anions or molecules.

Bearing solvent effects in mind, we first examined the reaction of the dienophile 5 with the diene 6 in solvents with diverse dielectric constants. As seen from Table 3, there is

Table 3. Homogeneous DA reactions catalyzed by the catalyst H2 in different solvents at room temperature^a

Entry	Solvent	Dielectric constant ²²	ee $(\%)$	Config.
	Et ₂ O	4.27	O	
2	THF	7.52	21	S
3	CH ₂ Cl ₂	8.93	15	S
	MeOH	33.00	40	S
	MeCN	36.64	35	S
6	MeNO ₂	37.27	24	

^a The reaction time was 22 h under the conditions the same as those given in [Table 1.](#page-189-0)

a correlation between the ees and the dielectric constants. Higher dielectric constants generally favor higher ees. Unlike the reaction of ethyl glyoxylate with 1,3-cyclo-hexadiene,^{[16a](#page-194-0)} there was no reversal in the product configuration. This indicates that the reversal we observed with H2 and S2 is less likely to be attributed solely to the change in the solvating media on going from homogeneous solution to solid surface. Because IR measurements of S2 indicate the presence of hydrogen bondings between the triflate anions and surface silanol groups, we suggest that the reversal could simply be resulted from the dissociation of the triflate ions from Cu(II) to the surface. This hypothesis implies that the triflate is coordinated to the PhBOX-Cu(II) complexes in homogeneous reactions. However, whilst there is evidence of one triflate coordination to Cu(II) at the solid state as well as suggestion of the association of triflate with Cu(II) in catalysis,^{[3c,23](#page-193-0)} the literature has generally assumed that the PhBOX-Cu(II)-dienophile intermediates adopt a distorted square planar configuration with O Tf^{$-$} playing little role.

To shed more light on the hypothesis that the reversal we observed resulting from the dissociation of OTf^- from the PhBOX-Cu(II) catalyst, we investigated the DA reaction catalyzed by the analogous $BOX-Zn(II)$ and $BOX-Mg(II)$ complexes derived from the corresponding triflates. In the absence of a coordinating ligand or counterion, the BOX- $M(II)$ -dienophile $(M=Zn, Mg)$ intermediates assume a tetrahedral geometry because of the lack of crystal field stabilization.^{[20](#page-194-0)} However, the homogeneous DA reactions catalyzed by the triflate complexes of BOX-Zn(II) and BOX- $Mg(II)$ are consistent with OTf⁻ coordination, forming octahedral BOX-M(II)-dienophile intermediates.^{18,20a} Thus, if immobilization leads to the removal of O^T from BOX-M(II) onto the surface, the heterogeneous BOX-M(II) catalysts would furnish products of opposite configuration. Table 4 summarizes the results we obtained with the catalysts H3, H4 and the corresponding immobilized catalysts S3 and S4. It is interesting to find that the

Table 4. Comparison of the immobilized catalysts S3 and S4 with the homogeneous analogues H3 and H4 for the DA reactions, respectively^a

Entry			Catalyst Solvent Conv. $(\%)$ <i>endolexo</i> ee $(\%)$			Config.
1	H3	CH_2Cl_2 > 97		89:11	22.	
2	S3	CH ₂ Cl ₂	95	85:15	24	
3	H4	CH ₂ Cl ₂	95	75:15	60	
$\overline{4}$	S4	CH ₂ Cl ₂	46	85:15	30	

^a The reaction time was 22 h under the conditions the same as those given in [Table 1](#page-189-0).

immobilization of H3 and H4 did result in the reversal of enantioselectivity, providing strong support for our hypothesis above that the reversal in product configuration between $H2$ and $S2$ is a result of $\text{O}\text{T}f^-$ dissociation from Cu(II) due to hydrogen bonding with the surface silanols. The lower activity observed with S4 is due to the lower loading of magnesium in comparison with that of zinc (0.03 vs $1.0 \text{ wt\%}.$

If anion coordination and dissociation is the cause for the product configuration reversal on going from H2–H4 to S2–S4, a similar reversal is also expected when the triflate counterion of H2–H4 is replaced with a non- or much less-coordinating anion. This is indeed the case with the PhBOX-Zn(II) and PhBOX-Mg(II) catalysts, as shown in Table 5. Thus, whilst the DA reaction catalyzed by H3 affords the S endo adduct with 22% ee (entry 1) which is consistent with a octahedral Zn(II) with axial triflate coordination, the product obtained by the (S)-PhBOX- $Zn(SbF₆)₂$ complex has the R absolute configuration with 64% ee (entry 2). In the latter case, the SbF_6^- counterion is believed to be fully dissociated, giving rise to a tetrahedral $Zn(II).^{20c}$ $Zn(II).^{20c}$ $Zn(II).^{20c}$ Likewise, with less coordinating perchlorate ion, (R)-PhBOX-Mg(II) furnished the product with configuration opposite to that observed with the triflate complex (Table 5, entries 3 and 4).^{[18](#page-194-0)}

In the case of the (S) -tBuBOX catalyst H1, the S endo enantiomer is always obtained regardless of the counterion, for example, SbF_6^- (non-coordinating) or OTf^- (coordina-ting).^{[3c](#page-193-0)} Thus, on the basis of the above analysis, the catalyst $S1$, in which O Tf⁻ dissociates due to hydrogen bonding, should give the same (S) *endo* enantiomer as the homogeneous H1 does. The results obtained are fully in accordance with this prediction ([Table 1,](#page-189-0) entries 1 and 2).

Applying the same argument to the (S) -PhBOX-Cu (II) complex, we were surprised to find that replacing the coordinating OTf⁻ anion with SbF_6^- and ClO_4^- did not result in the expected configuration reversal and in the case of (S)-PhBOX-Cu(SbF₆)₂, a *racemic* product was obtained (Table 5, entries 5, 6 and 8). The results obtained with (S) -PhBOX-Cu(ClO₄)₂ could be accounted for perchlorate coordination, leading to a square pyramidal or octahedral Cu(II) species and hence an adduct with S configuration. Although the 21-electron octahedral Cu(II) species not expected to be stable, there are a number of 5- and 6-coordinated Cu(II) complexes involving coordinated

^a The reactions were run in CH₂Cl₂ with other conditions the same as those given in [Table 1](#page-189-0). ^b The results were reported by Evans et al.^{3c}

^c The reactions were performed at -15 °C and were reported by Desimoni et al.^{[18c](#page-194-0)}

perchlorate ions.[24](#page-194-0) However this does not explain the loss of enantioface discrimination with (S) -PhBOX-Cu (SbF_6) .

Enlightened by the work of Jørgensen and co-workers,^{[16a](#page-194-0)} and aiming to gain further insight into the observations made above and particularly into the configuration reversal encountered with S2, we undertook HF (Hartree–Fork) modeling of the cationic BOX-M(II)-5 intermediates. For simplicity, the influence of the counter ions or the silanols was not considered during the calculation. The tBuBOX- $Cu(II)$ -5 and PhBOX-Zn(II)-5 are believed to prefer square planar and tetrahedral geometry, respectively, as aforementioned. Figure 5 shows the total energies of the two cations alongside that of PhBOX-Cu(II)-5 against the dihedral angle θ . As clearly seen, the Zn(II)-dienophile intermediate is indeed highly in favor of a tetrahedral geometry with the most stable configuration at θ = 80. The energy cost on going to the perfect square planar coordination is 12.4 kcal mol $^{-1}$ and to the tetrahedral arrangement is only 0.4 kcal mol^{-1}. In contrast, the most stable configuration for tBuBOX-Cu(II)-5 is found at $\theta = 39^\circ$, and it is significantly less energy-costly to change to the perfect square planar than to the tetrahedral geometry, 4.7 versus 9.3 kcal mol^{-1}. Thus the results from modeling confirm the speculations aforementioned, that is, the dienophile intermediate of tBuBOX-Cu(II) preferring square planar, and that of PhBOX-Zn(II) preferring tetrahedral in the absence of coordinating anions.

Figure 5. Calculated intermediate energies: (a) tBuBOX-Cu(II)-dienophile. (b) PhBOX-Cu(II)-dienophile, and (c) PhBOX-Zn(II)-dienophile. The dihedral angel θ refers to the angle between the plane of N–M–N and that of O–M–O.

Unlike those two intermediates, (S) -PhBOX-Cu(II)-5 has the most stable configuration at θ = 46°, right in the middle of a square planar and tetrahedral arrangement. More interestingly, the energy difference between going from the most stable state to the extreme planar and tetrahedral geometry is small, only about 1.7 kcal mol^{-1}. The energy needed for changing the dihedral angle from 46 to 80° , the most stable configuration for the tetrahedral PhBOX-Zn(II)- 5 which yields the R endo adduct, is 3.9 kcal mol^{-1}. This is only 0.5 kcal mol⁻¹ difference from that required for a change to the square planar geometry. Such a small energy difference makes us believe that (S) -PhBOX-Cu(II)-5 is flexible in configuration and is involved in a dynamic equilibrium between planar and tetrahedral geometries. This explains why the cyclization leads to a racemic product when the anion is non-coordinating and points to the enantioselection observed with the triflate salt of PhBOX-Cu(II) in the homogeneous DA reaction being a result of O Tf⁻ coordination to the PhBOX-Cu(II)-dienophile species. When 2 equiv H_2O was added to the reaction catalyzed by the $Sb\bar{F}_6^-$ salt, the S endo adduct was obtained albeit with a low 10% ee [\(Table 5,](#page-191-0) entry 7). This is consistent with water coordination, forming square pyramidal or octahedral Cu(II) intermediates and hence giving rise to the addition at the α -Si face of 5. This may also explain the effect of support on the ees discussed earlier ([Table 2,](#page-190-0) entries 7 and 8).

In the case of $S2$, in which evidence suggests that the $\text{OTf}^$ is dissociated, the DA reaction of 5 with cyclopentadiene 6 furnished the R endo adduct in 42% ee in toluene ([Table 2](#page-190-0), entry 4) instead of a racemic product as expected. This could be due to weak interactions between the surface Lewis acidic groups and the oxazolidinone ring oxygen of the substrate, and/or restrictions imposed by the surface on the conformations adoptable by the two phenyl rings. The formation of the R adduct is less likely to be purely a 'solvent' effect for an interfacial reaction, as solvents of widely different polarity did lead to the same face selection, although Jørgensen et al. found that face selection in the DA reaction of ethyl glyoxylate and 1,3-cyclohexadiene could be altered by solvents of different dielectric properties.^{[16a](#page-194-0)}

3. Conclusions

A series of the silica-supported heterogeneous BOX-M(II) catalysts have been successfully prepared for the DA reaction of $3-(E)$ -2-butenoyl)-1,3-oxazolin-2-one and cyclopentadiene at room temperature. The supported BOX-Cu(II) catalysts show enhanced enantioselectivities in toluene compared to their homogeneous counterparts, and it can be recycled without losing enantioselectivity. Spectroscopic evidence suggests that the immobilization of the homogeneous catalysts results from the hydrogen-bonding interactions between the triflate counterions and the surface silanol groups. Thus, as also shown by others, $11,12$ hydrogen bonding can provide a simple way for the immobilization of homogeneous catalysts, which requires neither modification of the catalysts nor functionalization of the surface.

One of the most interesting observations of this study is the reversal in product configuration when the homogeneous catalyst (S) -PhBOX-Cu $(OTf)_2$ is immobilized. Both the experimental results and theoretical calculations indicate that the triflate counterion coordinates to Cu(II) in homogeneous reactions and the configuration reversal upon immobilization is triggered by the dissociation of the anion from the metal cation due to hydrogen-bonding interactions with the surface silanols. Since a geometric change is highly feasible of the PhBOX-Cu(II)-dienophile intermediate towards either square planar or tetrahedral configuration and a conformational rearrangement of the phenyl rings of BOX is not expected to be energy-costly, the face selection and hence the enantioselectivity can be subtly affected by a number of factors such as the surface Lewis acidic and basic groups and the restrictions imposed by the surface morphology on the conformations adoptable by the BOX ligands.

4. Experimental

4.1. General

The 1 H NMR spectra were recorded on a 400 MHz spectrometer with $CDCl₃$ as the solvent. Elemental analyses were carried out on an inductively coupled plasma emission spectrometer (ICP-AES). FTIR spectra of the catalysts $(4000-400 \text{ cm}^{-1})$ were recorded on a Thermo Nicolet Impact 470 FTIR spectrometer. Self-supporting wafers of 1.3 cm were placed in an IR cell with $CaF₂$ windows. The wafers were purged at 60° C for 2 h under a controlled nitrogen atmosphere with a flow rate of 60 ml min⁻¹. The spectra were obtained by scans of 64 with a resolution of 4 cm^{-1} . Theoretical calculations were performed using the Gaussian 03 program package and were carried out at the HF (Hartree–Fork) Level with 3-21G* basis set for C, O, N, H and Lanl2dz basis set for Cu and Zn. All the optimized models were based on the known crystal structures.^{3c,16a} The solvent was not included in the calculation model.

4.1.1. Preparation of the heterogeneous catalysts. A $BOX-M(OTf)$ ₂ complex was dissolved in dichloromethane $(0.28 \text{ mmol}, \sim 0.03 \text{ M})$, and the resulting solution was filtered onto the pretreated silica (530 mg) under an argon atmosphere. The suspension was stirred for 3 h at room temperature, filtered, washed with dichloromethane several times and dried in vacuum to remove the solvent.

4.1.2. Heterogeneous DA reactions. An appropriate solution of dienophile 5 (0.23 mmol, 0.23 M) was added to a supported BOX-M(II) catalyst (metal content: 0.023 mmol), and the suspension was stirred for 15 min. 12 equiv of cyclopentadiene (2.8 mmol, 182 mg) was then added. After a certain period of time the reaction was stopped, and the catalyst was separated by filtration, thoroughly washed with the same solvent and then used for the subsequent reactions. The product was isolated by filtration through silica. The conversion was determined by ¹H NMR. The endolexo ratio and ee value of the endo isomer were analyzed by chiral HPLC [Chiralcel-OD column with hexane/ethanol (98/2) as the eluant]. The configuration of the products was confirmed by comparing with the literature. 36

4.1.3. Homogeneous DA reactions. An appropriate solution of dienophile 5 (0.4 mmol, \sim 0.23 M) was added to a solution of BOX-M(II) catalyst (0.04 mmol, \sim 0.03 M). The mixture was stirred at room temperature for 15 min, and

then 12 equiv of cyclopentadiene (4.8 mmol, 317 mg) was added. After a certain period of time, the reaction was stopped and the product was isolated by filtration through silica. The conversion, *endolexo* ratio and ee of the *endo* isomer were determined as described above.

4.2. Materials

2,2'-Isopropylidenebis[(4S)-4-tert-butyl-2-oxazoline] (tBu- $BOX)$ and $(S)-2,2'$ -isopropylidenebis(4-phenyl-2-oxazoline) (PhBOX) were purchased from Aldrich. $Cu(OTf)_{2}$, $Zn(OTf)_2$ and $Mg(OTf)_2$ were purchased from Fluka. Amorphous silica (pore diameter: 9.7 nm, BET: 390 m² g^{-1}) was commercially obtained. 3-((E)-2-Butenoyl)-1,3-oxazolin-2-one and the homogeneous BOX-M(II) catalysts were prepared following the literature procedures.^{3b,25} Commercially available compounds were used without further purification. Solvents were dried according to standard procedures. Amorphous silica was ground, washed with $1 M HNO₃$ and then with distilled water to neutrality, and dried in vacuum at 80° C. Before use, the silica was treated in air at $300\degree$ C for 3 h to remove the physisorbed water.

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- 3. Smith, D. H.; Masinter, L. M.; Sridharan, N. S. In Heuristic DENDRAL: Analysis of Molecular Structure; Wipke, W. T.; Heller, S. R.; Feldmann, R. J.; Hyde, E., Eds. Computer representation and manipulation of chemical information. John Wiley: New York, 1974; pp 287–298.
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